Pathologic myopia is a major cause of visual impairment worldwide. Pathologic myopia is distinctly different from high myopia. High myopia is a high degree of myopic refractive error, whereas pathologic myopia is defined by a presence of typical complications in the fundus (posterior staphyloma or myopic maculopathy equal to or more serious than diffuse choroidal atrophy). Pathologic myopia often occurs in eyes with high myopia, however its complications especially posterior staphyloma can also occur in eyes without high myopia.

Owing to a recent advance in ocular imaging, an objective and accurate diagnosis of pathologic myopia has become possible. Especially, optical coherence tomography has revealed novel lesions like dome-shaped macula and myopic traction maculopathy. Wide-field optical coherence tomography has succeeded in visualizing the entire extent of large staphylomas. The effectiveness of new therapies for complications have been shown, such as anti-VEGF therapies for myopic macular neovascularization and vitreoretinal surgery for myopic traction maculopathy.

Myopia, especially childhood myopia, has been increasing rapidly in the world. In parallel with an increase in myopia, the prevalence of high myopia has also been increasing. However, it remains unclear whether or not pathologic myopia will increase in parallel with an increase of myopia itself. In addition, it has remained unclear whether genes responsible for pathologic myopia are the same as those for myopia in general, or whether pathologic myopia is genetically different from other myopia.

Keywords: pathologic myopia, myopic maculopathy, optical coherence tomography, gene analyses, myopic macular neovascularization, myopic foveoschisis, myopic traction maculopathy, dome-shaped macula

**Definition of Pathologic Myopia**

According to the IMI, pathologic myopia is an excessive axial elongation associated with myopia that leads to structural changes in the posterior segment of the eye (including posterior staphyloma, myopic maculopathy, and high myopia-associated optic neuropathy) and that can lead to loss of best-corrected visual acuity. The term “pathologic myopia” is often confused with “high myopia.” These two entities are distinctly different; high myopia is defined as a high degree of myopic refractive error, whereas “pathologic myopia” is characterized by the presence of typical myopic lesions in the posterior fundus. Duke-Elder defined pathologic myopia as “that type of myopia which is accompanied by degenerative changes occurring especially in the posterior pole of the globe.” Characteristic myopic lesions in the posterior fundus include myopic maculopathy equal to or more serious than...
diffuse chorioretinal atrophy (equal to category 2 in the META-analysis for Pathologic Myopia [META-PM] classification) and/or the presence of a posterior staphyloma. The cut-off values of the myopic refractive error and axial length were not set for the definition of pathologic myopia because a posterior staphyloma has been reported to occur in eyes with normal axial length or in eyes with an axial length of less than 26.5 mm, although long axial length is one of the risk factors for fundus complications.

Because progressive choroidal thinning and formation of Bruch's membrane defects in the macular region are key phenomena associated with myopic maculopathy, the lesions of myopic maculopathy are better classified by their appearance in the optical coherence tomography (OCT) images than by the morphology as detected on fundus photographs.

**Basic Aspects of Pathologic Myopia**

Epidemiology of Pathologic Myopia

**Prevalence and Its Impact on Vision.** It has been reported that pathologic myopia affects up to 3% of the world's population, with racial differences regarding the prevalence of the disease. Approximately 1% to 3% of Asians and 1% of Caucasians have pathologic myopia. However, the definition of pathologic myopia used in earlier studies was not consistent and pathologic myopia was confused with high myopia. Thus, an accurate prevalence needs to be determined based on the new standard definition of pathologic myopia.

Pathologic myopia causes vision impairment or blindness in 0.2% to 1.5% of Asians and 0.1% to 0.5% of Caucasians. Particularly, it is one of the major causes of low vision in working-aged populations, as well as in the elderly population. In Asia, pathologic myopia is the leading cause of irreversible blindness in Taiwan, Japan, and China. In Taiwan, pathologic myopia is the second leading cause of vision impairment in individuals aged 65 years or older. In Japan, pathologic myopia is the third leading cause of bilateral low vision and the leading cause of monocular blindness in individuals aged 40 years or older. In China, pathologic myopia is the leading cause of blindness and low vision in individuals aged 40 to 49 years. In Western countries, pathologic myopia is the third cause of blindness according to the Rotterdam Study, the Copenhagen City Eye Study, and the Los Angeles Latino Eye Study.

**High Myopia and Myopia.** Recently, owing to changes in environmental and lifestyle factors, the prevalence of myopia and high myopia has increased rapidly. Therefore, the associated prevalence of pathologic myopia may also increase dramatically in the near future.

High myopia can be defined as a refractive error of at least −5.00 diopters (D). High myopia is linked to pathologic myopia. Most pathologic myopia occurs in eyes with high myopia, although low myopia and some individuals with emmetropia will also develop pathologic myopia. Approximately 28.7%, 44.4%, 45.9%, 47.6%, 58.31%, 72.7%, and 65% of high myopia cases in adult or elderly populations have pathologic myopia in Singapore, Australia, Japan, rural China, South China, Taiwan, and Beijing, respectively. This means that approximately one-half of subjects with high myopia in the adult population would develop pathologic myopia.

The prevalence of high myopia is linked to the overall prevalence of myopia. In Taiwan, the prevalence of myopia is as high as 84% in 18-year-old individuals and the prevalence of high myopia is about 21%. Approximately 22.9% of the world population were diagnosed with myopia in 2000, and 11.6% had high myopia among the myopia population. It is estimated that by 2050, approximately one-half of the world's population will have myopia and up to one-fifth of the myopia population will be highly myopic. Thus, areas with high prevalence rates of myopia would have increased rates of high myopia and pathologic myopia. Pathologic myopia will be associated with a high number of individuals suffering from vision impairment and will have a significant negative impact on society.

The degree of myopia is associated with the risk of pathologic myopia. The prevalence of pathologic myopia is only 1% to 19% in the low-to-moderate myopia (−3 D) population, but its prevalence is as high as 50% to 70% in the high myopia population. A 1-D increase in myopia is associated with a 67% increase in the prevalence of pathologic myopia. A linear trend is observed for increasing myopia dipters until −7.00 D, followed by an exponential trend in the prevalence rates of pathologic myopia.

**Age.** In addition to the degree of myopia, age is an important factor in the development of pathologic myopia. The prevalence of myopic maculopathy increased and exponentially with increasing spherical equivalent and age in Singapore. The prevalence of pathologic myopia is low in children and adolescents, but increases with advancing age. In individuals with high myopia aged 40 years or older, a progressive increase was noted in the prevalence and severity of maculopathy. Pathologic myopia changes with chorioretinal atrophy were found in 52.9% and 19.3% of the Chinese and Singaporean adult groups with high myopia, respectively. In the Taiwanese elderly population, myopic maculopathy increased from 2.2% in individuals aged 65 years to 14.8% in relatively older adults. Myopic maculopathy is rare in children with high myopia. However, a long-term follow-up study showed that 83% of adults with pathologic myopia and myopic maculopathy had already had diffuse choroidal atrophy around the optic disc in their childhood. This finding suggested the possibility that children who eventually develop pathologic myopia may be different, even at an early age.

**Genetic or Acquired.** There are two types of myopia—congenital myopia or infantile myopia and acquired myopia or school myopia. Congenital myopia has family aggregation and is strongly influenced by genetic factors. However, the prevalence of congenital myopia is low, less than 1% among Caucasian population. The initial degree of infantile myopia is often high and progression of myopia is also observed. A longer life span with high myopia may be linked to the high prevalence of pathologic myopia.

Acquired or school myopia occurs in children who develop myopia in the primary school or early secondary school years, with the exclusion of congenital myopia of strong familial inheritance. It should be noted that, from age 6, the annual progression rate for children with myopia is approximately 1 D until the end of adolescence, with the development of high myopia between the ages of 11 and 13 years. Considering the myopia boom in children worldwide, the severity of pathologic myopia associated with vision impairment is predictable.
Environmental Factors. In the case of acquired myopia, there might be a genetic susceptibility owing to a variation in the prevalence observed in different areas. Although more than 200 gene variants have been found to be associated with myopia, none predominately contribute to acquired myopia.\textsuperscript{35} Therefore, environmental factors play a more important role in the development of myopia and high myopia. Risk factors include educational stress, near-work time and intensity, and lack of outdoor time.\textsuperscript{36-37} In addition, increased digital screen time because of the smartphone era and the popularity of online education owing to the coronavirus disease 2019 pandemic may aggravate the prevalence of myopia in the near future.\textsuperscript{38}

Genetics

Genome-wide association studies (GWASs) have identified more than hundreds susceptibility genes for myopia.\textsuperscript{39-44} However, the genetic background for pathologic myopia has not been elucidated fully. For example, little is known about whether all subjects with high myopia have the same risk of developing pathologic myopia or if the risk of developing pathologic myopia depends on the patient's genetic background.

Candidate Gene Analysis for Pathologic Myopia.

Several studies have examined the association between myopic macular neovascularization (MNV) and susceptibility genes for myopia, high myopia, and AMD. Among the susceptibility genes for AMD and their related genes, ARMS2, CFH, C2/CFB, C3, CFI, ABCA1, APOE, LIPC, CETP, TIMP3, COL10A1, COL1A1, VEGFA, and PEDF were evaluated for their association with MNV.\textsuperscript{45-50} CFI, COL8A1, and PEDF were suggested as susceptibility genes for MNV,\textsuperscript{48-50} and it has been reported that VEGFA is associated with the size of MNV and the visual prognosis after treatment for MNV.\textsuperscript{51} However, these associations were not confirmed in later studies. Among the susceptibility genes for myopia and high myopia, CCDC102B, CCDC102A, COL8A1, RASGRF1, TOX, RDH5, and SHISA6 were evaluated for association with MNV, but no association was found.\textsuperscript{52,53}

In 2015, a photographic classification and grading system for myopic maculopathy was proposed.\textsuperscript{5} Since then, the Myopic Maculopathy Classification System (META-PM classification) has been used to study pathologic myopia. In 2019, the genotype distribution of 50 susceptibility genes for myopia were compared between 348 highly myopic cases with myopic maculopathy and 898 highly myopic controls without myopic maculopathy, but none of the genes showed a significant association with myopic maculopathy in the highly myopic eyes.\textsuperscript{54}

GWAS for Pathologic Myopia.

In 2018, a GWAS on myopic maculopathy in a Japanese population identified CCDC102B as a susceptibility gene for myopic maculopathy.\textsuperscript{55} The genotype distribution of CCDC102B was significantly different between the 1381 highly myopic cases with myopic maculopathy and the 936 highly myopic controls without myopic maculopathy. In contrast, CCDC102B was not significantly associated with axial length, and previous GWASs on myopia have not reported any association between CCDC102B and myopia. CCDC102B is a susceptibility gene for myopic maculopathy, but not for myopia. Given that the genetic background for developing myopia and the genetic background for developing myopic maculopathy are different, we would be able to develop preventive methods for myopic maculopathy, even for patients who have already developed myopia or high myopia. The role of CCDC102B in the development of myopic maculopathy should be elucidated.

The discovery of CCDC102B suggests that we might be able to prevent the development of myopic maculopathy, even after the development of high myopia. To control the development of pathologic myopia in highly myopic patients, further studies need to discover more susceptibility genes for myopic maculopathy that are not associated with high myopia. Because posterior staphyloma is associated with myopic maculopathy in highly myopic eyes, the identification of susceptibility genes for posterior staphyloma and/or posterior eye shape would also contribute to future control of the development of pathologic myopia. The posterior fundus shape can be quantitatively evaluated using OCT.\textsuperscript{56} GWASs on fundus shape might be able to discover genes associated with fundus shape and/or staphyloma and contribute to future control of the development of pathologic myopia.

Animal Models of Pathologic Myopia

Since Wiesel and Raviola\textsuperscript{57} discovered in 1977 that lid-sutured monkeys developed axial elongation, animal models have been a vital tool to understand and develop treatments for myopia. This section summarizes animals that spontaneously, or via monocular deprivation, developed features of pathologic myopia.

A search on Embase database using the following keywords: “animal model” with “posterior staphyloma,” “lacquer cracks,” “myopic maculopathy,” “choroidal neovascularization,” and “Fuchs spot” one feature at a time revealed 1502 results. There were 1466 entries from “choroidal neovascularization,” which were almost entirely induced artificially through laser photocoagulation and, hence, were excluded. A summary of screened studies is found in Table 1, which included retinopathy, globe enlarged (rge) chicks, normal chicks, and LRP2 knockout mice.

The murine model is widely used because its genetics and physiology are well-understood.\textsuperscript{58} Similar to in humans, murine sclera and fibroblasts contain five types of muscarinic receptors.\textsuperscript{59,60} However, they are nocturnal animals that lack a fovea and ability to accommodate.\textsuperscript{61} Additionally, their lenses are more spherical compared with human lenses and occupy most of the vitreal cavity.\textsuperscript{62}

Chick eyes are relatively larger and make up 50% of the cranial volume compared with 5% in adult human.\textsuperscript{63} Additionally, they contain pecten, a comb-like structure of blood vessels that provides nutrition and oxygenation to the retina. Crucially, chick sclera has a cartilaginous layer and ossicles, unlike any mammals.\textsuperscript{64,65} Nonetheless, there is a rod-free region called the area centralis in the chick retina that contains a greater concentration of cone photoreceptors, resembling the macula in human.\textsuperscript{66} Similar to mice, chicks have no fovea.

Animal Models of Lacquer Cracks.

Lacquer cracks are the yellowish linear lesions in the macular region and seem to lack choriocapillaris under OCT angiography (OCTA). They are seen in 4.3% to 15.7% of patients with pathologic myopia.\textsuperscript{67-70} These lesions are believed to represent mechanical breaks in the Bruch’s membrane–RPE–choriocapillaris complex. In pathologic myopia, lacquer cracks are commonly found temporal to the fovea in a horizontal direction.\textsuperscript{71} One may discuss that because a horizontal tension within Bruch’s membrane may be relieved
### Table 1. Summary of Animal Models That Demonstrated Features of Pathologic Myopia Spontaneously or Induced Through Monocular Deprivation

<table>
<thead>
<tr>
<th>Authors</th>
<th>Animal</th>
<th>n</th>
<th>Relevant Phenotypes</th>
<th>Method</th>
<th>Time of First Intervention</th>
<th>Degree of Induced Myopia</th>
<th>Other Features</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cases et al.,95 2015</td>
<td>Mouse</td>
<td>20</td>
<td>Peripapillary staphyloma after 21 days, chorioretinal atrophy after 60 days</td>
<td>Spontaneous in LRP2 knockout model. Mutation associated with Stickler syndrome and Donnai-Barrow or facio-acoustico-renal syndrome (DBS/FOAR), which present with ocular defect</td>
<td>Day 0</td>
<td>Day 90: About 40% increase in axial length</td>
<td>Increase in vitreous chamber, retina thinning (owing to increased cell death), scleral thinning at the posterior pole, and collagen fibrils formed fewer lamellae</td>
</tr>
<tr>
<td>Wen et al.,295 2006</td>
<td>Chick</td>
<td>45</td>
<td>Lacquer cracks after 12 weeks</td>
<td>Form-deprivation via lid suturing</td>
<td>Day 2</td>
<td>Week 4: −10.23 ± 2.15 D, Week 8: −15.57 ± 2.52 D, Week 12: −17.01 ± 3.29 D Axial length increases by 13.5%, 11.4% and 11.1% respectively</td>
<td>—</td>
</tr>
<tr>
<td>Hirata and Negi,72 1998</td>
<td>Chick</td>
<td>40</td>
<td>Lacquer cracks after 8 weeks, chorioretinal atrophy (choriocapillaris significantly less dense) after 4 weeks</td>
<td>Form deprivation via lid suturing</td>
<td>Day 1</td>
<td>Week 8: 28.3% increase in axial length</td>
<td>—</td>
</tr>
<tr>
<td>Montiani-Ferreira et al.,76 2004</td>
<td>Chick</td>
<td>9 rge/rge, 12 rge/+ , 5 +/+</td>
<td>Lacquer cracks after 6.4 weeks, patchy chorioretinal atrophy and scleral displacement after 134 weeks</td>
<td>Spontaneous in congenital stationnary night blindness model (mutation in GNB3 gene), animals known as retinopathy globe enlarged (rge) chicks</td>
<td>Day 0</td>
<td>Hyperopia. At Day 92, rge/rge shows 9.8 ± 7.4 D, rge/+ shows 4.3 ± 0.57 D. Axial length increases by 14.8%</td>
<td>Areas of ruptured Bruch’s membrane with focal absence of RPE Fibroblasts covered the interface of the abnormal Bruch’s membrane and choriocapillaris</td>
</tr>
<tr>
<td>Mao et al.,296 2006</td>
<td>Chick</td>
<td>32</td>
<td>Chorioretinal atrophy after 12 weeks</td>
<td>Form deprivation via lid suturing</td>
<td>Day 1</td>
<td>Week 12: −18.0 ± 2.25 D, 25.1% increase in axial length</td>
<td>TUNEL-positive cells in ONL and INL of myopic chick eyes</td>
</tr>
</tbody>
</table>

INL, inner nuclear layer; ONL, outer nuclear layer.
by a temporally located parapapillary gamma zone, lacquer cracks may be orientated in a horizontal direction to relieve the remaining excess tension in the vertical direction.

_Lid-sutured Chick_. Chicks that undergo monocular deprivation through lid suturing on the first day after hatching develop lesions that resemble lacquer cracks after 8 weeks. Similar to human, these lesions are frequently orientated horizontally, which may be due to the vertically orientated pecten in its retina. Vascular casts show that lacquer cracks are not sharply demarcated. There is a central gap that is void of Bruch’s membrane and choriocapillaris. This gap is surrounded by an area with reduced choriocapillaris density, possibly owing to atrophy from increased tension on the vessel bed. Large choroidal vessels bridging the gap could also be seen. Proliferation and accumulation of RPE, seen as a white arrow in Figure 1C, suggest that this lacquer crack has developed over a period of time and the RPE had sufficient time to respond.

_Retinopathy, Globe Enlarged (rge) Chick_. Rge chicks have a progressive, early onset visual loss and a deletion in GNB3, a gene that encodes guanine nucleotide binding protein b3 (Gbeta3), resulting in a nonfunctional protein. Physiologically, Gbeta3 forms a part of phototransduction cascade and is an integral part of the G-protein signaling within ON-bipolar cells. In humans, GNB3 mutation is seen in patients with congenital stationary night blindness with normal retinal thickness.

Lacquer cracks in rge chicks are seen as early as 45 days after hatching (Fig. 2A). Histologic images show a rupture of Bruch’s membrane and a lack of RPE compared with the surrounding area where these structures are intact (Fig. 2B).

**Long-term Follow-up Observation of Lacquer Cracks in Humans and Chicks.** Studies show that lacquer cracks in patients with pathologic myopia tend to deteriorate by elongating, increasing in number, or progressing to patchy atrophy. Interestingly, the chick models also develop similar progression patterns. In Figure 3, a 50-week-old lid-sutured chick eye developed a greater number of lacquer cracks and a marked elongation as compared with an 8-week-old eye shown in Figure 2. Although there was no obvious formation of a patchy atrophy during the long-term observation of lid-sutured chicks, 134-week-old rge chicks developed circular lesions that resembled a patchy atrophy (Fig. 5). OCT images at the lesions showed areas of tissue thinning and posterior bowing of sclera, which were consistent with a patchy chorioretinal atrophy in patients with pathologic myopia. This finding suggests that both features

---

**Figure 1.** Images taken from 8-week-old lid-sutured chick. (A) Macroscopic appearance of the eyecup of a lid-sutured eye. Horizontal lacquer cracks could be seen running perpendicularly to the pecten. (B) Vascular cast at site of the LC and showing a central gap surrounded by areas of decreased choriocapillaris density and some large choroidal bridging vessels between the gap. (C) Histology image at the site of the LC. The white arrow depicts the proliferation and accumulation of RPE while the red arrows indicates both ends of Bruch’s membrane.

FIGURE 3. Images of older lid-sutured (A) and rge chicks (B–D). (A) Eyecup of 50-week-old lid-sutured chick. There are more orientations and a higher number of lacquer cracks compared with an 8-week-old lid-sutured eyecup with two lesions crisscrossing each other (red arrow) (B) Confocal scanning laser ophthalmoscopy (cSLO) FA of 134-week-old rge chicks showing three circular lesions near the pecten. (C) Spectral domain OCT (SD-OCT) image of a circular lesion illustrating the marked loss of tissue and the posterior bowing of sclera at the lesion site. (D) A 3D image of the circular lesion showing similar findings. B, C, D reprinted with permission from Vutipongsatorn K, Nagaoka N, Yokoi T, et al. Correlations between experimental myopia models and human pathologic myopia. Retina. 2019;39:621–635.

may share a similar pathophysiology and that rge chicks might be a suitable animal model to study the disease.

Animal Models of Posterior Staphyloma. Posterior staphyloma is a localized, outpouching of the eye wall with a radius of curvature that is less than the surrounding curvature of the eye wall. It is present in approximately 50% of patients with pathologic myopia and little is known about its etiology. Scleral weakness secondary to tissue loss and change in collagen fibrils have been implicated in some studies. However, scleral re-enforcement treatments have yielded mixed results. Choroidal thinning has also been proposed because it is strongly associated with posterior staphylomas. However, patients with an acquired choroidal thinning still demonstrate axial elongation without developing posterior staphylomas. Furthermore, patients with retinitis pigmentosa can develop posterior staphylomas in areas where the choroid was intact. Bruch’s membrane is an acellular, five-layered extracellular matrix located between the RPE and choroid. With a thickness of 2 to 4 μm, it facilitates the transport of nutrients and metabolic byproducts between the choriocapillaris and the RPE, as well as providing structural support to the globe. Bruch’s membrane and its surrounding structures are markedly thinned or absent along the staphyloma edges. Additionally, eyes with a secondary Bruch’s membrane defect, such as in toxoplasmotic macular scars, can show a collateral scleral staphyloma. Furthermore, because its thickness remains constant while other layers thin out in axially elongated globes, Jonas et al. proposed that Bruch’s membrane may play an active role in the process of emmetropization, myopization and potentially staphyloma development as well.

Rge Chicks. As seen in Figure 3C, the OCT images of 134-week-old rge chicks show an outward bowing of the sclera in the regions of a Bruch’s membrane defect. Because the extent of scleral displacement is limited, it is difficult to conclude if these scleral deformations were certainly staphylomas. However, it seems reasonable to suggest that this may be an early stage of a transition toward a staphyloma formation. A longer observation period may be necessary to monitor the development of scleral deformation.
**LRP2 Knockout Mouse.** Currently, there is one animal model that definitively develops posterior staphyloma, which is the LRP2 knockout mouse. LRP2 (low-density lipoprotein receptor–related protein 2) encodes megalin—a transmembrane receptor found on the apical side of absorptive epithelia. It regulates the level of certain circulating compounds such as lipoproteins, sterols, vitamin-binding proteins, and hormones. Its mutation is associated with rare genetic conditions such as Donnai–Barrow syndrome and Stickler syndrome, both of which present with facial dysmorphology, hearing loss, ocular defects and chorioretinal atrophy.

Histologic sections and magnetic resonance imaging (MRI) show evidence of axial elongation and posterior staphyloma from days 15 and 21, respectively (Figs. 4A and 4B). A separation of the choroid and RPE began at the edge of the staphyloma. Both layers became progressively thinner before disappearing at the optic nerve head. Bruch’s membrane could not be identified clearly on these images. Furthermore, there was a marked thinning of the retina and pycnotic cell bodies were detected throughout the excavated region. This finding indicated an area of peripapillary atrophy apparently corresponding with the region with peripapillary staphyloma in this model. This formation is compatible

**FIGURE 4.** Images taken from 134-week-old rge chicks. (A) Confocal scanning laser ophthalmoscopy (cSLO) FA showing three circular lesions near the pecten. (B) Spectral domain OCT (SD-OCT) image of a circular lesion illustrating the marked loss of tissue and the posterior bowing of sclera at the lesion site. (C) A 3D image of the circular lesion showing similar findings.
CLINICAL ASPECTS OF PATHOLOGIC MYOPIA

Posterior Staphyloma

Posterior staphylomas are hallmarks of pathologic myopia and are among other major causes or sequel of developing myopic maculopathy. The presence of a posterior staphyloma is part of a recently updated definition of pathologic myopia that was characterized by the occurrence of myopic choroidal atrophy equal to or more serious than diffuse choroidal atrophy or by the presence of a posterior staphyloma.

As described by Spaide, a posterior staphyloma is an outpouching of a circumscribed posterior fundus region and has a curvature radius that is smaller than the curvature radius of the adjacent eye wall (Fig. 5). Posterior staphylomas should be differentiated from a simple scleral backward bowing which is commonly seen on OCT images of highly myopic eyes.

Applying three-dimensional (3D)-MRI, Moriyama et al. recently analyzed the shape of the whole eye, so that even large posterior staphylomas could be imaged entirely (Fig. 6). Based on 3D-MRI images of the eye, they showed that the difference in the ocular shape is correlated with the development of vision-threatening conditions in eyes with pathologic myopia. Applying 3D-MRI and wide-field fundus imaging, Ohno-Matsui recently classified posterior staphylomas into six types: the wide macular type, the narrow macular type, the peripapillary type, the nasal type, the inferior type, and other configurations (Fig. 7). This classification was based on the previous categorization of staphylomas by Curtin into 10 types, among which the types I to V were primary staphylomas and the types VI to X were compound staphylomas. The most predominant staphyloma type was the wide, macular type (74% of eyes with staphyloma), followed by the narrow, macular type of staphyloma (14% of eyes with staphyloma). However, 3D-MRI was not feasible as a screening technique and, owing to a relatively low spatial resolution, subtle changes of shallow staphylomas were difficult to detect.

A new prototype of a wide-field swept source OCT system has recently been developed and uses not one but
Classification of staphyloma. A new classification of posterior staphyloma according to its location and extent. The staphyloma type is renamed according to its location and distribution. Type I → wide, macular staphyloma, Type II → narrow, macular staphyloma, Type III → peripapillary staphyloma, Type IV → nasal staphyloma, Type V → inferior staphyloma, Others → staphylomas other than type I to V. Reprinted with permission from Ohno-Matsui K. Proposed classification of posterior staphylomas based on analyses of eye shape by 3D-MRI. *Ophthalmology*. 2014;121:1798-1809. © 2014 American Academy of Ophthalmology. Published by Elsevier Inc.

Multiple scan lines and generates scan maps allowing the 3D reconstruction of posterior staphylomas in a region of interest of $23 \times 20$ mm and a depth of 5 mm. Shinohara et al. showed that wide-field OCT can provide tomographic images of posterior staphylomas in a resolution and size that have been unachievable so far and that may replace 3D-MRI in assessing posterior staphylomas. Using wide-field OCT, the edge of staphylomas showed consistent features with a gradual thinning of the choroid from the periphery toward the staphyloma edge and a gradual rethickening of the choroid from the staphyloma edge in direction to the posterior pole, accompanied by a change in the curvature radius of the sclera at the staphyloma edge. The staphylomatous region shows a posterior outpouching of the sclera nasal to the staphyloma edge. (B) Three-dimensionally reconstructed image shows the staphyloma edge clearly (outlined by arrowheads). Reprinted with permission from Shinohara K, Shimada N, Moriyama M, et al. Posterior staphylomas in pathologic myopia imaged by wide-field optical coherence tomography. *Invest Ophthalmol Vis Sci.* 2017;58:3750-3758. Licensed under a Creative Commons Attribution 4.0 International License (CC BY).

Myopic Choroidal Atrophy

Myopic maculopathy, also known as myopic macular degeneration, is a key feature of pathologic myopia. Curtin and Karlin first described five myopic fundus changes that are associated with an increase of axial length, including optic nerve crescent, chorioretinal atrophy, central pigment spot, lacquer cracks, and posterior staphyloma. Later, Avila et al. developed a grading system of myopic maculopathy on a scale of increasing severity from 0 to 5 as follows: M0, normal-appearing posterior pole; M1, choroidal pallor and tessellation; M2, M1 plus posterior staphyloma; M3, M2 plus lacquer cracks; M4, M3 plus focal areas of deep choroidal atrophy; and M5, large geographic areas of deep chorioretinal atrophy shown as “bare sclera.” In an atlas of pathologic myopia, Tokoro updated and organized the lesions of myopic maculopathy. Tokoro classified myopic macular lesions into four categories based on ophthalmoscopic findings: (1) tessellated fundus, (2) diffuse choroidal atrophy, (3) patchy choroidal atrophy, and (4) macular hemorrhage.
Lacquer cracks were included in the diffuse atrophy category and macular hemorrhage was subclassified into two types of lesions—myopic MNV and simple macular hemorrhage. Later, Hayashi et al.98 investigated the natural course of 806 eyes of 429 consecutive patients with high myopia (myopic refractive error of > 8 D or an axial length of ≥ 26.5 mm) who had follow-up for 5 to 32 years. Hayashi et al.98 made some modifications to Tokoro’s classification based on clinical impression; they categorized lacquer cracks and myopic MNV as independent lesions. This longitudinal observation noted that the three plus lesions were added to these categories and were included as “plus signs”: (1) lacquer cracks, (2) myopic MNV, and (3) Fuchs spots. The reason for separating these plus signs from the categories is that these three plus lesions affect or potentially affect central visual acuity and may develop from, or coexist, in eyes with any categories of the myopic maculopathy. Based on this classification, pathologic myopia is defined as equal or greater than myopic maculopathy category 2, or presence of “plus lesion,” or the presence of posterior staphyloma.

Pathologic myopia is defined as equal or greater than myopic maculopathy category 2, or presence of “plus lesion,” or the presence of posterior staphyloma.

Highly myopic patients with a tessellated fundus are significantly younger than the patients with other lesions of myopic maculopathy.7,98,99 Fang et al.77 and Xiao et al.51 both showed that highly myopic eyes with a tessellated fundus had less myopia and a shorter axial length than those in the category 2 or above. Tokoro100 reported that approximately 90% of eyes with only a tessellated fundus had an axial length of less than 26 mm. The proportion of tessellation decreases linearly with longer axial length and tessellation is not seen in eyes with an axial length of more than 31 mm. In other words, almost all eyes with axial length of more than 31 mm would have progressed to advanced myopic maculopathy (i.e., diffuse atrophy or patchy atrophy).

The META-PM Classification of Myopic Maculopathy. Recently an international panel of researchers in myopia reviewed previous studies and proposed a simplified, uniform classification system for pathologic myopia (Table 2). In this simplified system (the META-PM classification), myopic maculopathy lesions are categorized into five categories from no myopic retinal lesions (category 0), tessellated fundus only (category 1), diffuse chorioretinal atrophy (category 2), patchy chorioretinal atrophy (category 3), to macular atrophy (category 4). Three additional features were added to these categories and were included as “plus signs”: (1) lacquer cracks, (2) myopic MNV, and (3) Fuchs spot. The reason for separating these plus signs from the categories is that these three plus lesions affect or potentially affect central visual acuity and may develop from, or coexist, in eyes with any categories of the myopic maculopathy. Based on this classification, pathologic myopia is defined as equal to or greater than myopic maculopathy category 2, or presence of plus lesion, or the presence of a posterior staphyloma.

Features of Each Lesion of Myopic Maculopathy.

Tessellated (or Tigroid) Fundus (Category 1). Tessellated fundus is defined by the increased visibility of large choroid vessels owing to axial elongation (Fig. 9). Tessellation begins to develop around the optic disc, especially in the area between the optic disc and the central fovea. A tessellated fundus alone does not affect the central vision, unlike the other lesions of myopic maculopathy (Table 2). A tessellated fundus, along with the myopic conus, is one of the preliminary visible signs in eyes with myopia in general and often observed in children with high myopia.101 Wong et al.111 also reported that a tessellated fundus and peripapillary atrophy were the most common findings in highly myopic Chinese adolescents (aged 12–16 years).

Highly myopic patients with a tessellated fundus are significantly younger than the patients with other lesions of myopic maculopathy.7,27,98,112 Fang et al.77 and Xiao et al.51 both showed that highly myopic eyes with a tessellated fundus had less myopia and a shorter axial length than those in the category 2 or above. Tokoro100 reported that approximately 90% of eyes with only a tessellated fundus had an axial length of less than 26 mm. The proportion of tessellation decreases linearly with longer axial length and tessellation is not seen in eyes with an axial length of more than 31 mm. In other words, almost all eyes with axial length of more than 31 mm would have progressed to advanced myopic maculopathy (i.e., diffuse atrophy or patchy atrophy).

The mean (or median) subfoveal choroidal thickness in eyes with tessellated fundus in high myopia varies from 80 to 166 μm,7,112–114 and is decreased almost by one-half compared with those with no myopic maculopathy in high myopia.7,113 The distribution pattern of the choroidal thickness in eyes with tessellation was different with those in eyes with no maculopathy, but was similar to greater myopic maculopathy categories (e.g., diffuse atrophy and patchy atrophy). This finding suggested that the tessellation might be the first sign for myopic eyes to become pathologic.

Hayashi et al.98 showed that only 13.4% of eyes with a tessellated fundus showed a progression after a follow-up period of 5 to 32 years; 10.1% developed diffuse chorioretinal atrophy, 2.9% developed lacquer cracks, and 0.4% developed a MNV in the first time progression. In the population-based Beijing Eye Study, progression was observed in 19% eyes with tessellation with 10 years of follow-up.100 Another large series of 810 highly myopic eyes that were followed for more than 10 years (mean follow-up, 18 years) showed a higher progression rate (27%).77 Because the eyes with greater myopic maculopathy categories showed a greater progression rate, it is suggested that myopic maculopathy tends to progress more quickly after the tessellated fundus stage. A tessellated fundus might be a relatively stable condition, and highly myopic eyes might stay in this condition for a relatively long period.

Diffuse Choroidal Atrophy (Category 2). Diffuse choroidal atrophy is observed as an ill-defined yellowish

<table>
<thead>
<tr>
<th>META-PM Classification*</th>
<th>Visual Impairment</th>
<th>Pathologic Myopia</th>
</tr>
</thead>
<tbody>
<tr>
<td>Tessellated fundus (category 1)</td>
<td>None</td>
<td>–</td>
</tr>
<tr>
<td>Diffuse chorioretinal atrophy (category 2)</td>
<td>Mild</td>
<td>+</td>
</tr>
<tr>
<td>Patchy chorioretinal atrophy (category 3)</td>
<td>Parafoveal scotoma</td>
<td>+</td>
</tr>
<tr>
<td>Macular atrophy (category 4)</td>
<td>Central scotoma</td>
<td>+</td>
</tr>
<tr>
<td>Plus lesions</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Myopic MNV (including Fuchs’ spots)</td>
<td>Central scotoma, distorted vision</td>
<td>+</td>
</tr>
<tr>
<td>Lacquer cracks</td>
<td>Temporal scotoma owing to simple hemorrhage, distorted vision (in some cases)</td>
<td>+</td>
</tr>
</tbody>
</table>

FIGURE 9. Fundus photographs showing different types of myopic maculopathy. (A) Right fundus showing a tessellated fundus with an axial length of 28.1 mm in a 37-year-old woman. The best-corrected visual acuity (BCVA) is 1.2. (B) Right fundus showing PDCA with an axial length of 29.76 mm in a 45-year-old woman. The BCVA is 0.8. (C) Right fundus showing MDCA with an axial length of 31.77 mm in a 76-year-old man. The BCVA is 0.5. (D) Left fundus showing patchy atrophy (arrows) with an axial length of 30.46 mm in a 60-year-old woman. The BCVA is 0.9. (E) Left fundus showing myopic MNV-related macular atrophy with an axial length of 33.16 mm in a 74-year-old woman. The BCVA is 0.15. (F) Left fundus showing patchy atrophy-related macular atrophy with an axial length of 32.95 mm in a 61-year-old woman. The BCVA is 0.2.

lesion in the posterior fundus of highly myopic eyes. The lesion is not uniformly yellow, but shows a granular appearance. However, the fundus color may look different according to the degree of fundus pigmentation among races. Diffuse choroidal atrophy primarily appear around the optic disc and increases with age and finally covers the entire posterior pole.116 Thus, diffuse choroidal atrophy is subclassified to peripapillary diffuse choroidal atrophy (PDCA)30 and macular diffuse choroidal atrophy (MDCA).37 The frequency of diffuse atrophy increases with age as well as with an increase of axial length.26,115,116 Tokoro et al.115 found that diffuse atrophy usually occurred around the age 40 and was observed in about 50% to 40% of patients after age 40. Recently, Liu et al.37 adopted the META-PM classification to evaluate the frequency and distribution of the diffuse choroidal atrophy according to the age, axial length and the best-corrected visual acuity. In this large Chinese highly myopic cohort, the proportion of diffuse choroidal atrophy in age groups of 7 to 11, 12 to 18, 19 to 39, and more than 40 years old was 20.9%, 9.2%, 23.1%, and 52.9%, respectively. The incidence of diffuse choroidal atrophy increased with longer axial length, from 3.6% in eyes with an axial length of less than 26.50 mm to 62.8% in eyes with an axial length of 28.50 mm or greater. Fluorescein angiography (FA) revealed a mild hyperfluorescence owing to tissue staining in the late phase of the angiogram.109 On indocyanine green angiography (ICGA), a pronounced decrease of the choroidal capillaries and medium and large-size choroidal vessels can be seen in the area of diffuse atrophy. OCT shows a marked thinning of the choroid in the area of diffuse atrophy. The subfoveal choroidal thickness in eyes with macular diffuse atrophy is usually less than 100 μm and the mean choroidal thickness is 50 μm based on a clinic-based study.11 In most cases, the choroid is almost absent, although it is sporadically present large choroidal vessels. Larger choroidal blood vessels can be observed to protrude to the retina. However, even in the area where most of the choroidal layer is absent, the RPE layer and outer retina are present. It might explain the relatively preserved vision in eyes with diffuse atrophy. With the use of OCTA, choriocapillaris flow impairment can be detected, even though the visualization of the choroidal circulation remains a challenge for interpretation in eyes with pathologic myopia. The OCTA in eyes with diffuse atrophy shows the low-density choriocapillaris, with the presence of medium and large choroidal vessels.111,116 Although the choroid becomes thinned in eyes with a tessellated fundus, the degree of choroidal thinning is much more serious in eyes with diffuse atrophy and such disproportionate thinning of choroid compared with the surrounding tissue (RPE, outer retina, and sclera) might be a key phenomenon in diffuse atrophy as well as pathologic myopia. Diffuse choroidal atrophy can be seen as a grayish-white, well-defined atrophy.109 Owing to an absence of RPE and most of the choroid, the sclera can be observed through transparent retinal tissue. The median size of patchy atrophy was 1.75 mm², varying from 0.05 to 101.3 mm²37 with the diameter of less than one or more lobules of the choriocapillaris. Pigment clumping is observed within the area of patchy atrophy, especially along the margin of the atrophy or along the large choroidal vessels. Abruptly emerging vessels are commonly observed within or near the edge of patchy atrophy, especially for those with large size.116 Patchy atrophy was found in 10.5% of patients in a clinic-based Japanese highly myopic cohort.117 The percentage of patchy atrophy increases linearly with age and reaches 32.5% after age 60 years.115 The prevalence of patchy atrophy is 3.3% in eyes with an axial length from 27.0 to 27.9 mm, and exceeds 25% and 50% if the axial length is longer than 31 mm and 32 mm, respectively.115 With time, the patchy atrophy enlarges and coalesces with each other.77,118,119,120 FA as well as ICGA show a choroidal filling defect in the area of patchy atrophy suggesting that this lesion is a complete closure of choriocapillaris.109 Fundus autofluorescence shows hypoafluorescence with distinct border owing to a loss of RPE in the area of patchy atrophy. Using OCT, patchy atrophy is characterized by the lack of RPE and outer retina with loss of most of choroid. Thus, the inner retinal layers have direct contact with the inner scleral surface. Swept source OCT also showed that discontinuities of Bruch's membrane in the area of patchy atrophy.117,120 The RPE terminates outside of the margin of the macular Bruch membrane defect. Patchy atrophy could be regarded as a Bruch's membrane rupture, not solely an atrophy. Patchy atrophy is subclassified into three types: patchy atrophy that develops from lacquer cracks P(Lc); patchy atrophy that develops within the area of an advanced diffuse chorioretinal atrophy P(D); and patchy atrophy which can be seen along the border of the posterior staphyloma.98 The shape of patchy atrophy may help to differentiate these types, because P(D) is usually circular or elliptical and P(Lc) is often longitudinally oval. P(Lc) is considered an enlargement of Bruch's membrane defect owing to lacquer cracks, and P(D) might also represent a Bruch's membrane hole developing within the area of advanced stage of diffuse atrophy. Almost all eyes (95%) with patchy atrophy progressed after a mean follow-up of 18 years, in which an enlargement of the original patchy atrophy was found predominantly in 98% and new patchy atrophy was found in 47% followed by development of myopic MNV in 21.7% and patchy-related macular atrophy in 8.3%.77 Miere et al.122 also reported that all patchy atrophies had significantly enlarged over at least 12 months using quantitative measurement. Such high percentages of progression of eyes with patchy atrophy could be explained by the biomechanical properties of Bruch's membrane so that as soon as a defect is created, the Bruch's membrane defect would enlarge over time with ongoing axial elongation. However, it is uncommon for extrafoveal patchy atrophy to later involve the central fovea. This means that it is rare for patchy atrophy to cause the central vision loss although this lesion leads to a paracentral absolute scotoma owing to a loss of photoreceptors within the atrophic area (Table 3). Macular atrophy is a well-demarcated, grayish-white or whitish, atrophic lesion centered on the fovea. The imaging features are similar to those of patchy chorioretinal atrophy. The main difference between patchy atrophy and macular atrophy is its location relative to the central fovea. Based on the long-term follow-up observation, it is suggested that macular atrophy could be subclassified into MNV-related macular atrophy and patchy atrophy-related macular atrophy. MNV-related macular atrophy develops centered in the central fovea and enlarges toward the periphery, and patchy atrophy-related macular atrophy develops outside of the foveal area and enlarging, or coalescing with other patchy atrophies, into the foveal center.77 The differentiation is mainly based on its morphologic features or is assisted by the history of MNV. The
## Table 3. Summary of Phase III Clinical Trials Using Anti-VEGF Agents for Myopic Choroidal Neovascularization

<table>
<thead>
<tr>
<th>Study</th>
<th>Treatment Groups</th>
<th>No. of Eyes</th>
<th>Mean BCVA Change at Study Primary End Point</th>
<th>Mean BCVA Change at Study Final Visit</th>
<th>Mean No. of Anti-VEGF Injections Over Study Period</th>
</tr>
</thead>
<tbody>
<tr>
<td>RADIANCE</td>
<td>Ranibizumab 0.5 mg guided by BCVA stabilization</td>
<td>106</td>
<td>+10.5 letters at month 3</td>
<td>+13.8 letters at month 12 month</td>
<td>4.6 ranibizumab injections over 12 months</td>
</tr>
<tr>
<td></td>
<td>Ranibizumab 0.5 mg guided by disease activity</td>
<td>116</td>
<td>+10.6 letters at month 3</td>
<td>+14.4 letters at month 12 month</td>
<td>3.5 ranibizumab injections over 12 months</td>
</tr>
<tr>
<td></td>
<td>vPDT then eligible to add ranibizumab 0.5 mg after month 3</td>
<td>55</td>
<td>+2.2 letters at month 3</td>
<td>+9.3 letters at month 12</td>
<td>2.4 ranibizumab injections from month 3 to 12</td>
</tr>
<tr>
<td>BRILLANCE</td>
<td>Ranibizumab 0.5 mg guided by visual stabilization</td>
<td>182</td>
<td>+9.5 letters at month 3</td>
<td>+12.0 letters at month 12</td>
<td>4.6 ranibizumab injections over 12 months</td>
</tr>
<tr>
<td></td>
<td>Ranibizumab 0.5 mg guided by disease activity</td>
<td>184</td>
<td>+9.8 letters at month 3</td>
<td>+13.1 letters at month 12</td>
<td>3.0 ranibizumab injections over 12 months</td>
</tr>
<tr>
<td></td>
<td>vPDT then eligible to add ranibizumab after month 3</td>
<td>91</td>
<td>+4.5 letters at month 3</td>
<td>+10.3 letters at month 12</td>
<td>3.2 ranibizumab injections from month 3 to 12</td>
</tr>
<tr>
<td>MYRROR</td>
<td>Aflibercept 2 mg</td>
<td>91</td>
<td>+12.1 letters at week 24</td>
<td>+13.5 letters at week 48</td>
<td>4.2 aflibercept injections over 48 weeks</td>
</tr>
<tr>
<td></td>
<td>Sham followed by aflibercept 2 mg after week 24</td>
<td>31</td>
<td>−2.0 letters at week 24</td>
<td>+3.9 letters at week 48</td>
<td>3.0 aflibercept injections from week 24 to 48</td>
</tr>
<tr>
<td>SHINY</td>
<td>Conbercept 0.5 mg</td>
<td>132</td>
<td>+12.0 letters at month 3</td>
<td>+13.3 letters at month 9</td>
<td>4.8 conbercept injections over 9 months</td>
</tr>
<tr>
<td></td>
<td>Sham followed by conbercept 0.5 mg after month 3</td>
<td>44</td>
<td>+0.6 letters at month 3</td>
<td>+11.3 letters at month 9</td>
<td>3.6 conbercept injections from month 3 to 9</td>
</tr>
</tbody>
</table>

BCVA, best-corrected visual acuity.
majority of macular atrophy is an atrophic stage of MNV, with a very few percentages related to secondary foveal involvement by enlargement of lacquer atrophy.

During an 18-year follow-up, loss of best-corrected visual acuity was associated with the development of MNV, MNV-related macular atrophy, and enlargement of MNV-related macular atrophy. At the last visit, most of eyes with a best-corrected visual acuity of less than 0.1 (20/200) had macular atrophy.

Lacquer Cracks (Plus Lesion). Lacquer cracks can be detected as fine, irregular, yellow lines in and around the macula. They are considered to represent healed and mechanical breaks of the RPE, Bruch’s membrane, and the choriocapillaris complex. Multiple lacquer cracks can often be seen in branching and crisscrossing patterns. The prevalence of lacquer cracks ranged from 4.2% to 15.7% in highly myopic eyes in several cohorts. Lacquer cracks can develop at a relatively early age in patients with pathologic myopia. Klein and Curtin reported that the mean age of patients with lacquer cracks was 32 years with a range of 14 to 52 years. Previous studies also indicated that lacquer cracks occur most often in eyes with an axial length between 29.0 mm and 32.0 mm. Lacquer cracks are easily overlooked overlying the rupture. Lacquer cracks are easily overlooked under the section on OCT because they are too narrow to detect. However, hypoautofluorescence, which is due to the atrophied RPE, can be stained as a fibrovascular scar tissue filling the Bruch’s membrane. When lacquer cracks are newly formed, MNV might develop through the Bruch’s membrane detachment. Three plus signs in the META-PM classification are considered as neovascular components.

The diagnosis of lacquer cracks is made based on multimodal imaging. ICGA is considered the gold standard for lacquer crack detection, observed as linear hypofluorescence in the late phase. On FA, lacquer cracks show a consistent linear hyperfluorescence during the entire angiographic phase, a window defect owing to RPE atrophy overlaying the defects of Bruch’s membrane in the early phase and a staining of healed scar tissue filling the Bruch’s membrane defect in the late phase. Fundus autofluorescence shows hypautofluorescence, which is due to the atrophied RPE overlaying the rupture. Lacquer cracks are easily overlooked on OCT because they are too narrow to detect. However, when the lesions are within the OCT scans, lacquer cracks appear as discontinuities of the RPE and increased transmission into the deeper tissue beyond the RPE. OCT shows partial defect of choriocapillaris in the region of lacquer cracks.

Xu et al. reported that 53.7% of eyes with lacquer cracks progressed during a mean follow-up of 3.5 years. Three progression patterns were found: increase in number (43.9%), elongation (9.8%), and progression to patchy atrophy (14.6%). The most common pattern was an increase in number of lacquer cracks. New lacquer cracks tend to occur perpendicularly from the existing lacquer cracks (branching) or in parallel with the existing lacquer cracks. An elongation of an existing lacquer crack is also seen in a small percentage of eyes. Lacquer cracks increase their width and progress to patchy atrophy. In some eyes, this progression is not a uniform widening of a pre-existing lacquer cracks, but small circular areas of patchy atrophy first develop along the lines of lacquer cracks, and then these circular areas enlarge and fuse with each other. Although lacquer cracks are often observed in the vicinity of MNV, it is unusual for MNV to develop secondarily from the existing lacquer cracks. This finding suggests that the lacquer cracks that are seen as yellowish linear lesions represent healed scar tissue. When lacquer cracks are newly formed, MNV might develop through the Bruch’s membrane rupture. However, once the Bruch’s membrane rupture is healed with scar tissue, a secondary MNV rarely occurs.

It is rare for lacquer cracks to develop across the central fovea itself. Thus, lacquer cracks themselves do not generally impair the central vision; however, the subretinal bleeding that develops at the onset of the rupture of Bruch’s membrane could cause the impairment of central vision even after absorption of the hemorrhage (Table 2).

Myopic MNV (Plus Sign). MNV is a major cause of central vision impairment in pathologic myopia. It has been included as a plus sign in the META-PM classification. MNV included three phases: the active phase with proliferation of a fibrovascular membrane including MNV, exudation, and hemorrhage; the scar phase exemplified by a Fuchs spot; and the atrophic phase represented by MNV-related macular atrophy. Thus, Fuchs spots were not considered to be independent lesions and they were a scar phase of MNV. Further details are discussed in the section on Myopic MNV.

New Classification and Grading System. Although the META-PM classification is well-suited to identify various stages of myopic maculopathy, this classification is only based on fundus photographs that could lead to an accurate diagnosis of atrophic lesions because of the different visual presentations according to the degree of fundus pigmentation among races. In addition, other myopic macular lesions such as myopic traction maculopathy and dome-shaped macula were not included. Thus, an OCT-based classification has been developed. Further details are discussed under the section on OCT-based Classification of Myopic Maculopathy.

Recently, Ruiz-Medrano et al. also published a comprehensive review summarizing the main features of pathologic myopia and proposed a new classification system based on three key factors—atrophy, traction, and neovascularization—or the ATN classification system. This proposed classification system does not make any changes to the current atrophy classification (five categories in the META-PM classification). Tractional component is newly included containing five stages of inner or and outer foveoschisis, foveal detachment, macular hole, and retinal detachment. Three plus signs in the META-PM classification are considered as neovascular components.

Progression of Myopic Maculopathy. Based on the META-PM classification, Fang et al. conducted a retrospective case series study including 810 eyes of 432 highly myopic patients who had been followed for 10 or more years. After a mean follow-up of 18 years, the progression of myopic maculopathy was observed in 58.6% for all eyes, with 74.3% for eyes with pathologic myopia at baseline. The three most frequent progression patterns were (1) an extension of peripapillary diffuse atrophy to macular diffuse atrophy in diffuse atrophy, (2) an enlargement of the original atrophic lesion in patchy atrophy, and (3) the development of patchy atrophy in lacquer cracks. From two Chinese population-based longitudinal studies, the 10-year progression rate of myopic maculopathy was 35.5% in elderly Chinese (aged ≥40 years) (the Beijing Eye Study) and the 5-year progression rate was also 35.3% in rural Chinese adult population (aged ≥30 years) (the Handan Eye Study). In another large highly myopic Chinese cohort (the Zhongshan Ophthalmic Center-Brien Holden Vision Institute High Myopia Cohort Study), approximately 15% of 657 highly myopic eyes had progression of myopic maculopathy over 2 years. Older age, longer axial length, and the presence of posterior staphyloma are the main factors associated with the development and progression of myopic maculopathy.

A Scheme Depicting the Progression Patterns of Myopic Maculopathy (Fig. 10). First, the progression
from category 0 (no myopic maculopathy) to category 1 (fundus tessellation) was not associated with a decrease in visual acuity. Although tessellation is not considered as pathologic myopia, a remarkable thinning of the choroid begins with the appearance of tessellation, which is the first sign of the progression of myopic maculopathy. Second, diffuse atrophy (category 2) primarily occurs in the peripapillary region (PDCA) and eventually extends into the macula (MDCA). Third, the eyes with patchy atrophy have a hole in the macular Bruch’s membrane that either forms by an enlargement of lacquer cracks or develops in regions of advanced diffuse atrophy with a more vulnerable Bruch’s membrane. Fourth, both patchy atrophy and macular atrophy (MNV related and patchy related) tend to enlarge with time. Sixth, macular atrophy is almost always MNV related, although patchy-related macular atrophy can occasionally occur.

Conclusion. The characteristics of each lesion of myopic maculopathy have been clarified by multimodal imaging with advanced technique. The wide use of the META-PM classification system allows researchers to perform direct comparisons across studies and provides a common tool for clinical trials and epidemiologic studies. Further studies targeting the pathogenesis of myopic maculopathy will be of benefit to find an effective treatment and finally impede the progression of myopic maculopathy.

Myopic MNV

Pathogenesis of Myopic MNV. The pathogenesis of myopic MNV is not understood fully, and several theories, such as the mechanical theory and the heredodegenerative theory have been proposed to explain the development of myopic MNV. The background changes in an eye with pathologic myopia are believed to contribute to the pathogenesis of myopic MNV. These structural changes involve multiple layers of the eye, including the RPE, Bruch’s membrane, choriocapillaris, and choroid, as well as the sclera, and are mostly driven by axial elongation. These changes can be observed clinically as an increasing severity of myopic macular degeneration. Specific lesions particularly associated with myopic MNV include lacquer cracks, patchy atrophy, and large myopic conus. Marked thinning of the choroid and loss of large choroidal vessels suggest that impaired choroidal perfusion may contribute to the development of progressive atrophy in a myopic macula.

Lacquer cracks are present in many eyes with myopic MNV and have been proposed to be an important predisposing lesion. Lacquer cracks are believed to represent ruptures in Bruch’s membrane and therefore mechanical stretching has been proposed as a potential underlying factor. Alterations in the cytokine levels have also been described in eyes with pathologic myopia and may contribute to the pathogenesis of myopic MNV. In eyes with myopic MNV, an increased VEGF level has been found in the aqueous humor compared with eyes undergoing cataract surgery. There have also been a suggestion that genetic or hereditary factors may play a role in the development of myopic MNV. Single nucleotide polymorphism in the complement factor I gene has been associated with myopic MNV.

Diagnosis of Myopic MNV. The diagnosis of myopic MNV is based on clinical findings. Patients may present with blurring, scotoma, or distortion of vision. Upon ophtalmoscopy, features of pathologic myopia such as diffuse or patchy choroidal atrophy or myopic conus are usually present. The myopic MNV typically appears as a flat, small, greyish subretinal lesion beneath, or in close proximity to, the fovea with or without hemorrhage. SD-OCT is a useful screening tool because it is noninvasive and can be performed rapidly. On SD-OCT, myopic MNV presents as a highly reflective area contiguous above the RPE (type 2 MNV), usually with minimal subretinal fluid (SRF).

A clinical diagnosis of myopic MNV is usually confirmed by FA, which demonstrates the presence of the neovascularization, which appears as a well-defined hyperfluorescence in the early phase with leakage in the late phase in a classic MNV pattern. More recently, OCTA has been shown to detect myopic MNV noninvasively with high sensitivity and specificity. The key advantage of OCTA lies in its noninvasive nature, which allows repeated scans to be performed at each visit. As such, some centers now accept the diagnosis of myopic MNV to be made by either form of angiography. However, because OCTA is a relatively new technology, users need to be aware of limitations including various artefacts and segmentation error. A key limitation of OCTA is that activity cannot be assessed reliably based on OCTA alone. Interpreting OCTA together with structural OCT is therefore recommended to fully assess the presence, type, area, and activity of the MNV.

Assessment of Myopic MNV Activity. An accurate assessment of activity of MNV is important in determining when to start treatment and whether further treatment is no longer warranted. A demonstration of leakage in FA has been the gold standard for evaluating the activity of MNV. However, FA cannot be repeated frequently owing to its invasiveness. Increasingly, SD-OCT has overtaken FA as the main modality for assessing activity, particularly during follow-up. During the active phase, myopic MNV typically appears as a dome-shaped, hyperreflective elevation above the RPE with ill-defined borders. In addition, a lack of RPE coverage may also help to differentiate an active MNV from an inactive or scarred MNV. SD-OCT is extremely useful in assessing features of coexisting myopic tractional maculopathy, which may be exacerbated by the treatment of MNV by anti-VEGF therapy. Although OCTA may detect the area of neovascularization, it does not reflect the level of activity. Flow signal within the MNV often persists in the scar or atrophic phase of myopic MNV.

Differential Diagnoses. Differential diagnoses to consider for myopic MNV include a simple macular bleed in a highly myopic eye, which is often associated with lacquer cracks. Using FA, simple macular bleeding appears as blocked fluorescence only and high flow signal is absent in OCTA. On SD-OCT, simple bleeds appear as a projection of the hemorrhage along the Henle’s fiber layer. The late phase of ICGA is also useful to confirm the absence of MNV and for detecting coexisting lacquer cracks as linear hyperfluorescences. Several inflammatory conditions may present with signs that can be confused with myopic MNV. The most common conditions are acute multifocal choroiditis and panuveitis, as well as punctate inner choroidopathy with or without secondary MNV. In myopic individuals greater than 50 years of age, neovascular AMD may be confused with myopic MNV. Polypoidal lesions may be seen at the edge of staphyloma or in eyes with tilted disc syndrome. Idiopathic MNV is diagnosed by excluding other causes; that is, it is a diagnosis of exclusion. Finally, serous detachment with or without neovascularization may occur in eyes with a dome-shaped maculopathy.
**Treatment of Myopic MNV.** The natural history of myopic MNV is generally poor without treatment.\textsuperscript{149–151} In a 10-year follow-up study that evaluated the long-term visual outcome of myopic MNV without treatment, visual acuity was decreased significantly at 10 years, with the proportion of eyes having a visual acuity of 20/200 or less increasing from 29.6\% to 88.9\% and 96.3\% at 5 and 10 years, respectively.\textsuperscript{150} Therefore, the treatment of myopic MNV is warranted to prevent progressive visual loss.

Based on the Verteporfin in Photodynamic Therapy (VIP) Study,\textsuperscript{152} verteporfin photodynamic therapy (vPDT) became the first approved treatment for myopic MNV as vPDT-treated eyes had better mean best-corrected visual acuity compared with sham treatment. However, vPDT was unable to result in a gain in mean visual acuity at 2 years.\textsuperscript{153} Moreover, vPDT also resulted in a significantly more frequent development of chorioretinal atrophy and significantly worse visual acuity compared with intravitreal anti-VEGF therapy.\textsuperscript{154} Therefore in the era of anti-VEGF therapy, the standard-of-care treatment for myopic MNV is the use of intravitreal anti-VEGF therapy and vPDT is not recommended.\textsuperscript{155–157}

The efficacy and safety of intravitreal anti-VEGF therapy for the treatment of myopic MNV has been evaluated in a number of large, phase III, multicenter, randomized, controlled clinical trials, including RADIANCE.\textsuperscript{158,159} BRILLIANCE,\textsuperscript{160} MYRROR\textsuperscript{161} and SHINY (Table 3).\textsuperscript{162,163} Both the RADIANCE and BRILLIANCE studies evaluated the use of intravitreal ranibizumab 0.5 mg versus vPDT,\textsuperscript{158–160} whereas MYRROR and SHINY compared the use of intravitreal aflibercept 2 mg and intravitreal conbercept 0.5 mg versus sham treatment, respectively.\textsuperscript{161,162} Results from these randomized controlled trials have all demonstrated conclusively that intravitreal ranibizumab, aflibercept, and conbercept resulted in significant mean visual acuity gains in patients with myopic MNV at their primary end points with an excellent safety profile. These positive findings have led to approval of these anti-VEGF agents for the treatment of myopic MNV by various health authorities. The off-label use of intravitreal bevacizumab and ziv-aflibercept, which were originally designed to treat systemic neoplasia, have also been evaluated for treating myopic MNV and both agents have resulted in favorable visual acuity gains after treatment.\textsuperscript{164–166} However, robust clinical trial data in using intravitreal bevacizumab or ziv-aflibercept for myopic MNV are lacking. The use of these anti-VEGF agents for myopic MNV should therefore be limited to patients with a lack of access to the on-label approved anti-VEGF agents.

In comparison with anti-VEGF treatment for MNV owing to neovascular AMD, the treatment burden in using anti-VEGF therapy for myopic MNV is considerably lower. The recommended treatment strategy in using intravitreal anti-VEGF therapy for myopic MNV is with a single initial injection followed by as-needed injection with regular monitoring using SD-OCT to assess for disease activity.\textsuperscript{155,156} This strategy is based on the treatment protocols used in the ranibizumab disease activity-guided arms of the RADIANCE and BRILLIANCE studies\textsuperscript{158,160} and in the aflibercept arm of the MYRROR study.\textsuperscript{161} The effectiveness of this as-needed treatment approach has been evaluated in various real-world studies.\textsuperscript{168,169} The prospective LUMINOUS study demonstrated in the real-world setting that ranibizumab for myopic MNV resulted in a mean visual improvement of 9.7 letters and 1.5 letters at 1 year in treatment-naive and previously treated eyes, with a low mean number of ranibizumab injections of 3.0 and 2.6 injections over 12 months respectively.\textsuperscript{168} Another large-scale prospective real-world study evaluating the use of ranibizumab for myopic MNV in Japan also demonstrated similar findings, with a mean improvement in the logMAR best-corrected visual acuity of 0.19 unit and a low mean number of 2.0 injections over 12 months.\textsuperscript{169} The long-term results of the RADIANCE study also confirmed the favorable visual outcomes and low number of retreatment, with a mean visual acuity gain of 16.3 letters and 83\% of patients required no further treatment for myopic MNV after up to 48 months of follow-up.\textsuperscript{170}

Several studies have evaluated the prognostic factors associated with various treatment outcomes in using anti-VEGF therapy for myopic MNV.\textsuperscript{159,171,172} A subgroup analysis of the RADIANCE study showed that ranibizumab treatment resulted in a significant visual acuity gain, regardless of the level of baseline age, ethnicity, lesion area, MNV location, severity of myopia, axial length, and presence or absence of SRF.\textsuperscript{152} It was found that eyes with a larger baseline MNV lesion area required more injections over the study period compared with those with a smaller area.\textsuperscript{159} A post hoc analysis of eyes in the MYRROR study also showed that the severity of myopic macular degeneration severity did not seem to influence the visual or anatomical outcomes after intravitreal aflibercept treatment for myopic MNV.\textsuperscript{171} Another study showed that better visual acuity outcomes after ranibizumab or bevacizumab treatment might be associated with a shorter duration of symptoms, better baseline best-corrected visual acuity, and fundus autofluorescence pattern.\textsuperscript{172}

One of the main issues concerning the long-term visual outcomes of myopic MNV after anti-VEGF therapy is the development of myopic MNV-related macular atrophy, which can result in a gradual loss of the visual acuity initially gained over the long run.\textsuperscript{157,165} Because the current anti-VEGF therapy for myopic MNV only targets the angiogenesis process, future research efforts should therefore consider investigating methods to target macular atrophy associated with myopic MNV to achieve better long-term visual outcomes.

**Myopic Traction Maculopathy**

**Pathogenesis.** Myopic traction maculopathy includes a variety of pathologies secondary to tractional force on the retina in highly myopic eyes. Epiretinal membrane, lamellar hole, and many other conditions generating traction to the retina are included. Among myopic traction maculopathies, myopic foveoschisis is unique to pathological myopia and worth to learn as a specific complication. Because of the space limitation, this article will shed a light on this pathology. Myopic foveoschisis is referred to as a posterior retinal detachment without a macular hole in highly myopic eyes, first documented as a case report by Phillips in 1958.\textsuperscript{173} Almost 40 years later, Takano and Kishi\textsuperscript{174} found that retinoschisis and foveal detachment are common in highly myopic eyes in OCT. Myopic foveoschisis is characterized by various foveal architectural abnormalities, including a foveal cyst, a lamellar hole, and a foveal detachment.\textsuperscript{175}

Typical OCT images of this disease showing split between inner and outer retina within posterior staphyloma have led to the hypothesis that the inner retina is less flexible than the outer retina.\textsuperscript{179} Factors limiting the inner retinal flexibility include the vitreous cortex adhering to the retina, epiretinal membranes, internal limiting membrane (ILM), and retinal vessels. Preretinal membranes are often hard to recognize in high myopia, however it is present at the microscopic
FIGURE 11. Typical appearance of myopic foveoschisis. The fundus photograph (inset) shows a slightly elevated retina at the posterior pole, although it is not clearly identifiable. A horizontal OCT scan involving the macula shows retinoschisis in multiple retinal layers and a retinal detachment at the fovea (*asterisk*). There is glial tissue bridging the inner and outer layers of the retinoschisis (a so-called column, *arrow*).

Retinal detachment arising from a macular hole is a typical complication in highly myopic eyes. The vitreous cortex adhering to the retinal surface around the hole causes tangential traction that generates an inward vector component in deep staphyloma in highly myopic eyes, resulting in a retinal detachment.178

Assessment.

Symptoms. Patients are normally aware of central visual distortion corresponding with the involved area for retinoschisis and a relative scotoma for retinal detachment. Patients may be aware of an absolute scotoma at the center of the relative scotoma when a macular hole opens within a retinal detachment. Patients also report visual field loss at the involved area if an extensive retinal detachment is complicated. The Watzke–Allen test is usually negative for macular hole within the area of retinal detachment.

Fundus and OCT Appearance. Myopic foveoschisis can be recognized as a slight elevation of the posterior retina in highly myopic eyes; however, OCT is essential for accurate diagnosis especially in an atrophic fundus. OCT are essential not only for complete assessment of the retinal status but for surgical decision-making. Myopic foveoschisis presents with retinoschisis in multiple retinal layers (Fig. 11). The split retinal layers normally have a bridge between them, the so-called column, which is presumed to be residual Müller cells.179 ILM separation from the other retinal layers can also be observed, a so-called ILM detachment (Fig. 12A), and is a good indicator of the tractional force from the ILM.180 The tent-like peak of the inner retina is observed on the OCT image. This finding is coincident with retinal vessels and the so-called retinal microvascular traction, and more clearly observed after vitreous surgery with ILM peeling (Fig. 12B).181 This tractional force to the retinal vessels are also observed as a paravascular microhole in highly myopic eyes.182 The ellipsoid zone line of the photoreceptors sometimes disappears in the area of the retinal detachment;183 however, the ellipsoid zone line is typically well preserved in the area of retinoschisis. This finding suggests that the photoreceptor function is well-preserved in this subtype.

Based on OCT appearance, there are two stages before macular hole formation associated with retinoschisis (Fig. 13). The first stage is the development of the retinoschisis type, in which only retinoschisis is present and not a retinal detachment (Fig. 13A).176 Several months (sometimes several years) later, a retinal detachment begins around the fovea. This stage is the so-called foveal detachment type (Fig. 13B). After a while, the inner retina above the detachment is stretched and torn. This is how a macular hole appears as a consequence of retinoschisis with a retinal detachment.

There are two types of macular holes in highly myopic eyes. (Fig. 14)184 One is the type with the edge of the hole thickened with retinal cysts (Fig. 14A). There is no retinal detachment around the hole clinically, and this type usually does not change for months or years. The other has surrounding retinoschisis instead of retinal cysts around the hole (Fig. 14B). This type of macular hole results from myopic foveoschisis and typically progresses rapidly because of underlying traction.

Treatment.

Surgical Indications and Results. Investigators have reported that the vision decreased in 69% of patients, a macular hole developed in 31% after 3 years of follow-up,185 and in 50% of patients with retinoschisis a macular hole or retinal detachment developed after 2 years.186 These
IMI Pathologic Myopia

FIGURE 13. Time course and two distinct subtypes of myopic foveoschisis of the same patient shown in Fig 11. (A) The OCT image at the initial visit showing the retinoschisis type characterized by only retinoschisis without a retinal detachment. (B) One year later, the foveal detachment type occurs, which is characterized by a small localized retinal detachment (asterisk). The photoreceptors are separated from the RPE.

FIGURE 14. OCT appearance of two distinct subtypes in highly myopic macular holes. (A) A macular hole without retinoschisis has only retinal cysts and is normally stable, whereas (B) a macular hole with surrounding retinoschisis usually presents a higher likelihood of consequent retinal detachment.

observations encourage surgery on myopic foveoschisis to prevent more serious condition, namely, macular holes.

Asymptomatic myopic foveoschisis is not a good surgical indication because of a significant number of patients who suffer surgically induced visual decrease. The chance of visual improvement after surgery is much greater in cases with a foveal detachment than retinoschisis alone. The mean visual acuity after vitrectomy for macular hole with retinal detachment was less than 20/200, and the initial reattachment rate was about 50% to 70% after vitrectomy. Once a macular hole develops, the hole is difficult to close after vitrectomy in highly myopic eyes. The macular hole closure rate with retinoschisis or retinal detachment is about 40% to 50% based on OCT images. This difficulty probably arises because of the presence of posterior staphyloma for which the retina is extremely stretched. In fact, a longer axial length is generally a poor prognostic factor.

Vitrectomy. The vitreous plays an important role; therefore, creating a posterior vitreous detachment is critical. To create posterior vitreous separation, a vitreous cutter and silicone-tipped backflush needle with active suction are normally used. Because multiple components such as vitreous cortex, epiretinal membrane, and ILMs are tightly adhering to the retinal surface, a diamond-dusted membrane scraper facilitates to peel these extremely thin membranes. Importantly, great care is needed to separate the vitreous at the macula because of the risk of macular hole formation.

The necessity for ILM peeling remains controversial in myopic foveoschisis. However, the ILM is separated from the other retinal layers on OCT images in most eyes, suggesting that the rigidity of the ILM plays a significant role. ILM peeling also can enhance macular hole closure and remove any traction on the retina in myopic macular holes with a retinal detachment and raises the success rate. Indocyanine green or brilliant blue G are commonly used to stain the ILM selectively.

It is hypothesized that traumatic damage at the fovea induced by ILM peeling may cause full-thickness tissue loss at the fovea in myopic foveoschisis. This idea led to an attempt for surgeons to leave the foveal ILM. Recently, a nonfoveal ILM peeling technique was introduced to avoid macular holes. So far, studies reported a significant reduction of the incidence of postoperative macular holes; however, the visual acuity level was similar even with a higher anatomic success rate.

Macular holes in eyes with high myopia are much less likely to close. The inverted ILM flap technique was applied for highly myopic cases. The visual benefit has not been proven, especially in cases with retinal detachment; however, the macular hole closure rate seems to be higher than with the conventional ILM peeling technique. A study using this inverted ILM flap technique has shown a significantly higher success rate than conventional peeling technique. Autologous ILM or partial thickness or full-thickness retinal transplants have been reported recently to enhance the closure of myopic macular holes.

Because there is no coagulation maneuver around the macula, it is critical to support the retina for a long time to recover the integrity between the RPE and photoreceptors. It is preferred to use long-acting gas tamponade (i.e., perfluoropropane) to attain a greater anatomic success rate. Silicone oil tamponade is also an effective option.

Postoperative Complications. The opening of a macular hole occurs in between 10% and 20% of patients who underwent vitrectomy for myopic foveoschisis. Cases in which ILM peeling was performed in the presence of an extremely thin, stretched fovea are at risk of developing this complication. We investigated the incidence of postoperative macular hole formation and looked for the risk factors. A preoperative discontinuity of the ellipsoid zone line defect seen on
OCT was a predictor of macular hole development. Other factors, such as age, gender, and visual acuity, did not affect the rate of macular hole formation.

The recurrence of a retinal detachment is a major postoperative complication after vitrectomy for a macular hole with a retinal detachment. Removing residual vitreous cortex and epiretinal membranes is critical during a reoperation. However, it is sometimes difficult to identify the apparent cause of a re-detachment. In those cases, persistent traction, such as microvascular traction, might be responsible. Macular buckling can be considered in these cases. Macular buckling can compensate for any tractional force and showed a slightly higher initial reattachment rate than vitrectomy in eyes with a macular hole with a retinal detachment.\textsuperscript{205,206}

**Conclusion.** With the advancement in ocular imaging technologies, the pathophysiology of myopic macular diseases has become better understood. A number of interesting findings have suggested the presence of underlyling traction in highly myopic eyes. The preoperative features of these pathologies should be evaluated carefully, as these are important for decision-making and for assessing more effective surgical procedures in patients with myopic macular complications.

**Dome-Shaped Macula**

**Definition and Clinical Features.** In 2008, a French team reported a previously undescribed cause of visual loss in myopic patients that they have called dome-shaped macula.\textsuperscript{207} It was characterized by an inward bulge of the macula within the chorioretinal posterior concavity of the eye. This macular bulge was hardly detectable on fundus biomicroscopy, and was mainly visualized by OCT. They have reported five unilateral and five bilateral cases, corresponding with 15 of the 140 myopic eyes in their database. The degree of refractive changes ranged between –2 D and –15 D. A visual loss was found in 11 eyes, with a median visual acuity of 20/50. The authors have also reported various degrees of changes in the RPE, including atrophy and pigment clumping, observed in all eyes. A focal fluorescein leakage was found in seven eyes, but MNV could be ruled out in all cases.\textsuperscript{207} The finding has been confirmed by other authors in various countries and continents, allowing a better characterization of the condition. The prevalence of dome-shaped macula has been estimated at between 10.7% and 14.6% in the Chinese myopic population.\textsuperscript{208,209} Most reported cases are adult patients, but the condition has also been reported in children and adolescents.\textsuperscript{210} Mild-to-moderate myopia is usually reported, but the condition may also be observed in highly myopic eyes, in emmetropic eyes, and, less often, in hyperopic eyes.\textsuperscript{211,212} Subtypes of dome-shaped macula have been identified depending on the dome shape: round (Fig. 15) or oval with a vertical (Fig. 16) or horizontal axis (Figs. 17 and 18).\textsuperscript{213–215} In these studies, horizontal dome-shaped macula accounted for two-thirds to three-quarters of the cases. It is thus recommended to perform OCT B-scans along different axes to make the diagnosis of dome-shaped macula.

**Imaging.** In the first descriptions, time-domain OCT was used and it was unable to demonstrate if the bulge was due to the sclera or not.\textsuperscript{207,212} OCT has shown that the sclera was significantly thicker in the macular area in myopic eyes with dome-shaped macula compared with myopic eyes without dome-shaped macula,\textsuperscript{216} whereas there was no difference in the scleral thickness outside the macular area. Thus, the authors have concluded that dome-shaped macula could be the result of a relatively localized change in the scleral thickness beneath the macula in their highly myopic patients. In another study, dome-shaped macula has been found to correspond to a ridge between two outward concavities within the posterior staphyloma.\textsuperscript{217} Recently, 3D-MRI has allowed the reconstruction of the entire posterior pole in these eyes and has shown the presence of morphological changes in the entire posterior pole, with a hand-shaped inward convexity that extended horizontally from the optic disc through the fovea in most eyes with a horizontal oval dome-shaped macula, and a round inward convexity of the macular area in most eyes with a round dome-shaped macula. This MRI study has also shown a high diversity in convexities that could be the border of a single staphyloma or multiple staphyloma areas.\textsuperscript{209} The results from studies assessing the choroidal thickness are more controversial. A relatively thick choroid has been reported in some studies,\textsuperscript{211,212} whereas other studies have reported a relatively thin choroid,\textsuperscript{218} and a similar choroidal thickness has been found between eyes with and without dome-shaped macula with the same axial length in a comparative case study.\textsuperscript{219} However, the mean central choroidal thickness seemed to be thicker than the choroid in the surrounding staphyloma, and the mean central choroidal thickness-to-mean perimacular choroidal thickness ratio seemed to be greater in dome-shaped macula eyes than in other myopic eyes with the same axial length.\textsuperscript{220} Thus, a dome-shaped macula seems to be associated with a relatively thicker sclera and choroid at the bulge apex.

**Pathogenesis.** Dome-shaped macula has been suggested to be a possible compensatory mechanism in myopic anisometropia, after comparing the two eyes of a 49-year-old woman with mild myopic anisometropia.\textsuperscript{221} However, little is known about the occurrence of the condition, which may be unilateral or bilateral. Ohno-Matsui et al.\textsuperscript{212} have observed a peri dome choroidal deepening. Based on this finding, they have assumed that there might be a dome-induced inward push of the Bruch's membrane at the dome-shaped macula apex, leading to a compression of the subfoveal choroid. They have also observed Bruch's membrane defects in 20% of their studied eyes, suggesting the presence of Bruch's membrane strains on the dome's slopes.\textsuperscript{222}

**Course and Complications of a Dome-shaped Macula.** Long-term studies have shown that the bulge height increases moderately over time.\textsuperscript{218,219} This increased height has been shown to be associated with the extension of macular atrophic changes.\textsuperscript{219} The scleral thickness slightly decreases over time, both in the fovea and in parafoveal areas.\textsuperscript{218}

The main complication of dome-shaped macula is the occurrence of a macular serous retinal detachment (SRD). A macular SRD has been observed in all studies, but its incidence varies from 1.8% of cases in a Japanese cohort to 54% of cases in a French cumulative study.\textsuperscript{214,220} A dome-shaped macula is considered a vision-threatening condition, accounting for most cases of visual loss. A SRD is more commonly observed when the bulge height is greater than 350 microns and in eyes with vertical oval dome-shaped macula.\textsuperscript{213,223} The pathogenesis of SRD remains unknown but, it has recently been shown that the submacular choroidal blood flow measured on the OCTA B-scan is greater in eyes with SRF than in eyes without SRF.\textsuperscript{220} Thus, the changes in the choroidal thickness at the
Round dome-shaped macula (DSM). The diagnosis of DSM may be challenging based on fundus examination (A, —). The border of the staphyloma (yellow line) appears on the wide-field color photo (C). FA (B) only shows a slight hyperfluorescence in the macula. The DSM is visible on both SD-OCT horizontal (D) and vertical (E) B-scans, which also reveal a serous macular detachment. The choroid is relatively thick in the subfoveal area. The 3D reconstruction of the macula (F, G) clearly shows the round inward bulge of the macula.

transition between the bulge and the staphyloma observed by different teams\textsuperscript{17,224} may correspond with abrupt changes in choroidal blood flow, leading to outer blood-retina barrier failure and resulting in the occurrence of SRF. The spontaneous course of SRF is characterized by major changes in SRF height that may sometimes result in the complete resolution of the fluid.\textsuperscript{225} The fluid resolution may also correspond with the development of RPE and outer retina atrophy in the macula.\textsuperscript{219} Most studies of SRF associated with dome-shaped macula have shown a poorer visual acuity at baseline or during the follow-up\textsuperscript{219,226} but in one study, a similar prognosis has been reported between eyes with and without SRF after a 2-year follow-up.\textsuperscript{227}

Other reported complications include pigment epithelial detachment at the dome apex,\textsuperscript{228} polypoidal choroidal vasculopathy,\textsuperscript{229} and MNV that is more easily detected on OCTA.\textsuperscript{228,230} A similar response of MNV to ranibizumab has been reported between eyes with and without dome-shaped macula in a comparative series\textsuperscript{231} and in the RADIANCE trial.\textsuperscript{232}

Differential Diagnosis. Dome-shaped macula and its complications are very similar to the inferior staphyloma commonly observed in eyes with tilted disc syndrome. Indeed, both conditions include a change in the eyeball curvature\textsuperscript{233} that may lead to complications such as pigmentedary changes, MNV, polypoidal choroidal vasculopathy, and
Vertical dome-shaped macula (DSM). Fundus examination (A) is not contributive. Fundus autofluorescence (B) shows a hypoautofluorescence revealing RPE atrophy in the macula, surrounded by hyperautofluorescent areas. FA shows an uneven hyperfluorescence with leaking points (C). The difference in curvature is obvious when comparing the OCT horizontal (D) and vertical (E) B-scans, or on the 3D reconstruction of the posterior pole (F). A SRD is present. MRI shows a macular focal thickening of the eye wall in the right eye (G, arrow).

Similarities between these conditions have been highlighted in different studies, and similar hypotheses have been suggested to explain the occurrence of SRF. A condition close to dome-shaped macula called ridge-shaped macula has been recently described in younger patients (<20 years of age). It is characterized by a macular elevation in only one meridian across the fovea, observed in the horizontal direction across the vertical scan of OCT. In this study, no staphyloma and no Bruch’s membrane defect were associated.

Therapeutic Attempts. No treatment for dome-shaped macula is available currently. Its complications such as MNV or polypoidal choroidal vasculopathy should be treated promptly with anti-VEGF agents. Different therapies have been proposed for SRF complicating dome-shaped macula, including photodynamic therapy, spironolactone, aflibercept, and topical carbonic anhydrase inhibitors. A transient improvement has been observed after surgery for epiretinal membrane. However, all studies were case reports or small series and no comparison with a control group was performed. To date, no effective treatment is available for SRF complicating dome-shaped macula.

Glaucoma

The definition of pathologic myopia as suggested by the META-PM Study Group describes the morphological abnormalities in the macular region of highly myopic eyes, while coexisting changes in the optic nerve head have been mostly omitted. Optic nerve damage, however, relatively often coexists with maculopathy in eyes with pathologic myopia, and not quite rarely, it is an additional cause for visual field defects and vision impairment. In population-based studies of various ethnicities, the increase in the prevalence of glaucomatous or glaucoma-like optic neuropathy occurred beyond a myopic refractive error of –8 D or an...
axial length of more than 26.5 mm.\textsuperscript{248,250} It suggested that the prevalence of glaucoma in eyes with low to moderate levels of myopia or in eyes with an axial length shorter than 26.5 mm was similar to that for emmetropic eyes. The same values of a myopic refractive error of \(-8\) D or an axial length of 26.5 mm were the cut-off values for the definition of high myopia, when an enlargement of the optic disc and parapapillary regions was used as biomarker for the presence of high myopia.\textsuperscript{251,252}

For the diagnosis and for a better understanding of glaucoma in myopic eyes, the appearance of the optic nerve head is of great importance. The optic nerve head changes can be divided into those occurring in non–highly myopic eyes, and those in highly myopic eyes. In non–highly myopic eyes, the major change associated with axial elongation is the shifting of Bruch’s membrane opening in spatial relationship to the lamina cribrosa, usually in direction to the macula.\textsuperscript{253}

Considering the optic nerve head being composed of three layers, the Bruch’s membrane opening as the inner layer, the choroidal opening as the middle layer, and the opening in the peripapillary scleral flange with the lamina cribrosa as the outer layer, the Bruch’s membrane opening shift into the temporal direction leads to an overhanging of Bruch’s membrane into the intrapapillary region at the nasal optic disc margin, and to a lack of Bruch’s membrane in the temporal parapapillary region. It explains the development of the parapapillary gamma zone that is usually located in the temporal to temporal inferior parapapillary region and that is defined by the absence of Bruch’s membrane.\textsuperscript{254–258} A hypothetical cause for the Bruch’s membrane opening shift backward in direction to the macula may be a production of new Bruch’s membrane in the equatorial region for axial elongation in the process of emmetropization and myopization.\textsuperscript{93} As a sequel of the Bruch’s membrane opening...
shift, occurring usually in the horizontal plane, the optic disc shape, as assessed on ophthalmoscopy, changes from a mostly circular shape to a vertical oval form. Another reason for the ophthalmoscopic change to a vertical optic disc shape is a perspective distortion of the ophthalmoscopic image of the optic disc, owing to the lateralization of the optic disc. In non–highly myopic eyes, the size of the optic disc is not correlated significantly with the axial length of the refractive error. Correspondingly, the size of the Bruch’s membrane opening is not related to the axial length in non–highly myopic eyes.

In high myopia, the size of the Bruch’s membrane opening and of the optic disc (defined as the region with the lamina cribrosa as its bottom), and consequently the size of the lamina cribrosa, increase with longer axial length. The enlargement of the Bruch’s membrane opening leads to the development of a circular gamma zone, that is, the Bruch’s membrane opening enlargement is more marked than the enlargement of the optic disc (and lamina cribrosa) is. In association with the enlargement of the optic disc and lamina cribrosa, the latter gets elongated and thinned. The lamina cribrosa thinning decreases the distance between the intraocular compartment with the IOP and the retrobulbar compartment with the orbital cerebrospinal fluid pressure. Both pressures form the translaminar cribrosa pressure difference which is of high importance for glaucomatous optic neuropathy. The decreased distance between both compartments leads to a steepening of the translaminar cribrosa pressure gradient what may be one of the reasons for the increased glaucoma susceptibility in highly myopic eyes. Another reason for the increased glaucoma susceptibility may be morphological changes inside of the lamina cribrosa in association with its stretching.

In the parapapillary region, highly myopic eyes show, in addition to the gamma zone, the development and enlargement of the parapapillary delta zone. The delta zone is the equivalent of an elongated (stretched) and
thinned peripapillary scleral flange (within the gamma zone). The latter is the continuation of the inner portion of the posterior scleral into the lamina cribrosa, with the collagen fibers of the peripapillary border tissue of the scleral flange crisscrossing at the border of the scleral flange to the lamina cribrosa. The peripapillary scleral flange forms the anterior border of the orbital cerebrospinal fluid space and is covered only by the peripapillary retinal nerve fiber layer. It forms the biomechanical anchor for the lamina cribrosa. The outer portion of the posterior sclera merges with the optic nerve dura mater, with the circular arterial circle of Zinn–Haller usually located at the merging line. The arterial circle serves roughly as an ophthalmoscopic peripheral demarcation line of delta zone. The axial elongation-associated lengthening of the peripapillary scleral flange thus leads to an increased distance between the arterial circle and the lamina cribrosa that is nourished by the branches of the arterial circle. The increased distance between the arterial circle and the lamina cribrosa may be a further reason for an increased vulnerability of the optic nerve head in high myopia. Fitting with the role of the peripapillary scleral flange as the biomechanical anchor of the lamina cribrosa, a clinical study has suggested that a larger delta zone is associated with a higher prevalence of glaucomatous optic neuropathy in highly myopic eyes. In a parallel manner, a larger optic disc was correlated with a higher glaucoma susceptibility, whereas in contrast the size of gamma zone was not significantly related to the prevalence of glaucoma.

The importance of the peripapillary border tissues for the biomechanics of the optic nerve head and the glaucoma susceptibility has remained elusive so far. The peripapillary border tissue are differentiated into the peripapillary border tissue of the scleral flange (Elschnig) and the peripapillary border tissue of the choroid (Jacoby). The first one is the continuation of the optic nerve pia mater and, as mentioned elsewhere in this article, crisscrosses with the fibers of the scleral flange when the flange continues into the lamina cribrosa. The second one is interposed between the peripapillary border tissue of the scleral flange (Elschnig) and the end of Bruch’s membrane. It forms the border between the choroidal space and the intrapapillary prelaminar compartment. Because the inner shells of the eye, that is, the uvea, Bruch’s membrane, and the retina, are directly or indirectly connected to the outer ocular shell, that is, the sclera, only at the scleral spur anteriorly and through the peripapillary choroidal border tissue posteriorly, the latter may have biomechanical importance for the whole globe in general, and in particular for the connection between the inner ocular shells bearing the IOP and the lamina cribrosa. Owing to the development of the gamma zone, the distance between the end of Bruch’s membrane and the optic border elongates. It leads to a thinning of the peripapillary choroidal border tissue. It has been discussed that, in some highly myopic eyes, the choroidal border tissue may rupture, leading to a free Bruch’s membrane end and a corrugation of Bruch’s membrane in its vicinity. The peripapillary border tissues may, thus, be of some importance for the increased glaucoma susceptibility in high myopia.

In agreement with the histologic findings and with observations made in population-based investigations, a recent clinical study revealed that the prevalence of glaucomatous optic neuropathy increased from approximately 12.0% in eyes with an axial length of less than 26.5 mm to 28.5% in eyes with an axial length of 26.5 mm or greater and less than 28.0 mm, to 32.6% in eyes with an axial length of 28 mm or greater and less than 29 mm, to 36.0% in eyes with an axial length of 29 mm or greater and less than 30 mm, and to 42.1% in eyes with an axial length of 30 mm or greater. After adjusting for axial length, a higher glaucoma prevalence was correlated with a larger delta zone and a larger optic disc. These findings indicate that clinically a large optic disc and/or large delta zone should arouse the suspicion of a coexisting glaucomatous optic nerve damage in highly myopic eyes.

From a practical point of view, the detection of glaucomatous optic nerve damage gets more difficult with longer axial length. Axial elongation leads to a flattening of the physiological cup, so that the difference between the height of the neuroretinal rim and the bottom of the optic cup decreases, and makes the outlining of the cup/rim border more difficult. In addition, the pinkish color of the neuroretinal rim is reduced, so that the color contrast between the rim and cup is decreased. The increased brightness of the parapapillary region, mainly owing to the gamma zone and delta zone, decreased the detectability of the retinal nerve fiber layer, and the occurrence of both zones make the detection of glaucoma-associated beta zone difficult. Also, highly myopic eyes with glaucomatous optic neuropathy tend to have an IOP within the normal range, and perimetric defects are often unspecific, because they may be caused by refractive abnormalities, myopic macular changes, and/or nonglaucomatous optic nerve damage. The most important sign for the presence of glaucomatous optic nerve damage in highly myopic eyes may be the ophthalmoscopically detected intrapapillary vessel kinking relatively close to the disc border so that the inferior–superior–nasal–temporal rule of the shape of the neuroretinal rim is no longer fulfilled.

The detection of glaucoma progression as compared with the detection of the presence of glaucoma is even more difficult, in particular in highly myopic eyes. Observation of changes in the outer visual field may perhaps help for the detection of glaucoma progression since most other psychophysical and morphometric techniques may fail. Although glaucomatous optic neuropathy in highly myopic eyes can be defined by the intrapapillary vessel kinking relatively close to the optic disc border, studies are lacking that show the dependence of the effect of this optic nerve damage on the IOP. One may therefore speak of glaucomatous or glaucoma-like optic neuropathy. In addition, there may be a nonglaucomatous optic nerve damage in eyes with a large gamma zone on the temporal side. In a recent population-based study, a higher prevalence of high myopia or myopic maculopathy was correlated with a thinner retinal nerve fiber layer after exclusion of glaucomatous eyes. It fits with clinical observations that highly myopic eyes without myopic maculopathy category III or IV and without a glaucoma-like optic nerve head could show a deep central scotoma. One may discuss that the elongated distance between the disc and fovea, caused by the enlargement of gamma zone, led to a stretching of the macular retinal ganglion cell axons and to their loss. Such a mechanism may mimic glaucomatous optic nerve damage. It has remained unclear whether a myopic traction maculopathy, with the inner limiting membrane lifting the retinal ganglion cell layer and thus relatively shortening the distance between the retinal ganglion cells and the optic disc, may be partially protective against such a nonglaucomatous optic nerve damage in highly myopic eyes. If that...
OCT-Based Classification of Myopic Maculopathy

OCT has dramatically improved our understanding of various retinal pathologies in last two decades. Recent advances in OCT technology, including enhanced depth imaging OCT and swept source OCT, provided the high-resolution images and highly reliable measurements of choroid in highly myopic eyes. It has been well-known that the choroid in high myopia is markedly thin. Thus, adding the choroidal thinning of the choroid (almost absence of the entire choroid) in that diffuse atrophy was characterized by an extreme thinning of the choroid in highly myopic eyes. Earlier studies showed that choroidal attenuation may have a great contribution to the pathogenesis of myopic maculopathy. In addition, swept source OCT revealed that patchy atrophy and myopic MNV-related macular atrophy were not simply an atrophy, but were a macular Bruch’s membrane defect. Furthermore, OCT enables to detect new macular lesions which were not identified by fundus photographs, such as myopic traction maculopathy and dome-shaped macula.

Although the META-PM classification is very useful, the diagnosis of diffuse atrophy relies solely on its yellowish appearance on ophthalmoscopy. However, the fundus color may look different according to the degree of fundus pigmentation among races, which could impact the accurate diagnosis of atrophic lesions. Earlier studies showed that diffuse atrophy was characterized by an extreme thinning of the choroid (almost absence of the entire choroid) in OCT images. Thus, adding the choroidal thickness into the classification makes the diagnosis possible, objective, and more accurate. In addition, other myopic macular lesions that can only be diagnosed with OCT, such as myopic traction maculopathy and dome-shaped macula, can be included in the OCT-based classification of myopic maculopathy (Table 4).

Key OCT Features Determining Pathologic Myopia.

Extreme Choroidal thinning. It is well-known that the choroid in highly myopic eyes is thinner compared with normal eyes. In eyes with pathologic myopia (defined as myopic eyes equal to or more serious than diffuse atrophy), the thinning of the choroid is extreme. In many cases, almost the entire thickness of the choroid disappears with sporadically remaining large choroidal vessels.

A marked thinning of the choroid starts temporal to the optic disc in childhood, as PDCA. The presence of PDCA in children with high myopia is considered to be an indicator for the eventual development of advanced myopic maculopathy in later life. On OCT images, PDCA in children is characterized by a profound and abrupt thinning of the choroid in the temporal parapapillary region. The thinning of the choroid is abrupt and local; thus, it is critically different from a generalized impairment of choroidal circulation. Subsequently, Yokoi et al. also compared the OCT findings between a hospital-based group of highly myopic children with PDCA and a population-based control group of children. It was shown that 76% of the PDCA group had a choroidal thickness at 2500 μm nasal to the fovea that was thinner than 60 μm, which was not found in the control group. It indicated that this potential cut-off value of choroidal thickness may be helpful for the detection and diagnosis of PDCA in myopic children. With time, the area of extreme choroidal thinning enlarges and it is recognized as MDCA.

Bruch’s Membrane Holes. The defects of Bruch’s membrane in the macular region were reported in a histologic study of highly myopic eyes. Other macular lesions specific to pathologic myopia, that is, with myopic MNV-related macular atrophy and with patchy atrophy, had a choroidal thickness at 2500 μm nasal to the fovea which was not found in the normal fundus, whereas in eyes with myopic maculopathy, the subfoveal choroid was the thinnest. This observation was also confirmed in vertical scan by another study comparing choroidal thickness in eyes with tesselated and normal fundus. What is more, the distribution pattern...
**Table 4. OCT-based Classification of Myopic Maculopathy**

<table>
<thead>
<tr>
<th>Stage of MM</th>
<th>New Terminology</th>
<th>Details</th>
<th>Old Terminology</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ia</td>
<td>Peripapillary choroidal thinning</td>
<td>CT of &lt;56.5 μm at 3000 μm nasal from the fovea, outside of gamma zone.</td>
<td>PDCA</td>
</tr>
<tr>
<td>Ia Plus sign</td>
<td>Linear Bruch’s membrane defects</td>
<td>Yellowish linear lesions. Discontinuities of the RPE and increased hypertransmission into deeper tissues beyond the RPE in OCT image.</td>
<td>Lacquer cracks</td>
</tr>
<tr>
<td>Ib</td>
<td>Macular choroidal thinning</td>
<td>CT of &lt;62 μm subfoveally.</td>
<td>MDCA</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Plus sign Linear Bruch’s membrane defects</td>
<td></td>
</tr>
<tr>
<td>II</td>
<td>Extrafoveal Bruch’s membrane defects</td>
<td>Well-defined, grayish white, round lesion(s) in the macular, extrafoveal area. The Bruch’s membrane defect is usually surrounded by a slightly wider RPE defect the size and shape of which determines the ophthalmoscopic size and shape of the lesion. In the region of the Bruch’s membrane defect, the outer retinal layer, the RPE, the choriocapillaris and most of the medium-sized choroidal vessel layer are absent, and a medium-sized or large choroidal vessel may occasionally be present. The middle and inner retinal layers, more or less thinned, are in direct contact with the inner scleral surface</td>
<td>Patchy atrophy</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Plus sign Myopic CNV</td>
<td></td>
</tr>
<tr>
<td>III</td>
<td>Foveal Bruch’s membrane defect</td>
<td>The OCT-based histology is similar to the histology of extrafoveal Bruch’s membrane defects.</td>
<td>CNV</td>
</tr>
<tr>
<td>III Plus sign</td>
<td>Foveal Bruch’s membrane defect,</td>
<td></td>
<td>Macular atrophy</td>
</tr>
<tr>
<td></td>
<td>CNV-related</td>
<td>The OCT-based histology is similar to the histology of extrafoveal Bruch’s membrane defects.</td>
<td></td>
</tr>
<tr>
<td>IIIb</td>
<td>Foveal Bruch’s membrane defect,</td>
<td>Well-defined, round lesion including the fovea and expanding centrifugally around the fovea. The edges of the macular Bruch’s membrane defect are often upturned. In the center, remnants of Bruch’s membrane can be present, folded up in the process of the RPE-associated scar formation.</td>
<td>CNV-MA</td>
</tr>
<tr>
<td>IIIa Plus sign</td>
<td>Foveal Bruch’s membrane defects,</td>
<td></td>
<td>Patchy MA</td>
</tr>
<tr>
<td></td>
<td>patchy related</td>
<td>develops outside of the foveal area and enlarges in direction to the fovea or coalesces with other extrafoveal Bruch’s membrane defects in direction to the fovea.</td>
<td></td>
</tr>
<tr>
<td>Plus sign</td>
<td>Macular traction maculopathy</td>
<td>Schisis-like inner retinal splitting, schisis-like outer retina splitting, foveal detachment, lamellar or full-thickness macular hole and/or macular detachment.</td>
<td></td>
</tr>
<tr>
<td>Plus sign</td>
<td>Dome-shaped macula and macular ridge</td>
<td>Dome-like, inward bulge of the RPE line in all meridians with a height of &gt;50 um above a base line connecting the RPE lines on both side outside the dome. Macula ridge: Inward bulge of the RPE line in either horizontal or vertical scan with a height of &gt;50 um above a base line connecting the RPE lines.</td>
<td></td>
</tr>
</tbody>
</table>

CT, choroidal thickness; CNV, choroidal neovascularization; CNV-MA=choroidal neovascularization-related macular atrophy; Patchy MA, patchy atrophy-related macular atrophy.
FIGURE 19. Graph showing the pattern of distribution of choroidal thickness (CT) at different locations for each type of myopic maculopathy in the horizontal (A) and in the vertical direction (B). The CT was measured at the subfoveal region and at 3 mm nasal, temporal, superior, and inferior to the fovea. MA = macular atrophy.

FIGURE 20. The subfoveal CT (mean with standard deviation) in each myopic maculopathy are shown. The CT decreased significantly from normal fundus to tessellated fundus, to PDCA, and to MDCA in all locations. There is no significant difference in choroidal thickness between eyes with MDCA and patchy atrophy. *P < 0.05. CT = choroidal thickness; MA = macular atrophy; NS = not significant.

of choroidal thickness in eyes with a tessellation was similar to the eyes with other myopic maculopathy in greater categories, such as, PDCA, MDCA, and patchy atrophy. This finding suggested that the tessellation might be the first sign for myopic eyes to become pathologic.

Progressive Choroidal Thinning From Tessellation to PDCA, and to MDCA, But Not Thereafter. It was shown that the mean subfoveal choroidal thickness in eyes with a normal fundus was 274.5 μm, tessellated fundus was 129.1 μm, PDCA was 84.6 μm, MDCA was 50.2 μm, patchy atrophy was 48.6 μm, MNV-related macular atrophy was 27.3 μm, and patchy-related macular atrophy was 3.5 μm (Fig. 20). The choroidal thickness in all locations decreased from normal fundus to tessellated fundus, to PDCA, and to MDCA, as the severity of the myopic maculopathy increased. But there was no significant difference in choroidal thickness between eyes with MDCA and patchy atrophy in all locations except for those at nasally. Because the other two studies did not differentiate MDCA from PDCA, it is reasonable that the choroidal thickness in diffuse atrophy, which might include both PDCA and MDCA, was significantly thicker than those eyes with patchy atrophy.

The subfoveal choroidal thickness of eyes with macular atrophy (both MNV related and patchy related or either of them) was significantly thinner than that in any other groups. The subfoveal choroidal thickness in eyes with patchy-related macular atrophy was even thinner than choroidal thickness in MNV-related macular atrophy. It was noted that there was no difference among MNV-related macular atrophy, MDCA, and patchy atrophy in choroidal thickness at temporal, nasal, superior, and inferior locations.

The result from other two studies also indicated that the choroidal thickness was negatively correlated with severity of myopic maculopathy. A high myopia clinic-based study from Zhongshan Ophthalmic Center investigated the choroidal thickness in a large population. In this study, the median subfoveal choroidal thickness was 165 μm in normal fundus (C0), 80 μm in tessellated fundus (C1), 49 μm in diffuse atrophy (C2), 35 in patchy atrophy (C3), and 6.5 μm in macular atrophy (C4). It also showed that
the subfoveal choroidal thickness became significantly thinner with the increasing severity of maculopathy in C0 to C4 but the eyes with C3 to C4 shared similar parafoveal choroidal thickness, leading to the conclusion that C4 was not the result of progression from C3 which was consistent with long-term follow-up study by Fang et al.77

Cut-Off Value of Choroidal Thickness for Identifying PDCA and MDCA. Fang et al. used a receiver operating characteristic curve and Youden's index to determine the optimal cut-off choroidal thickness value of 56.5 μm in nasal choroidal thickness (3000 μm from fovea) for the diagnosis of PDCA. The performance is particularly excellent in the less than 20 years of age group with a high sensitivity of 90% and a good specificity of 88%. The area under curve of choroidal thickness in each location became lower in the older age group. For the 60 to 79 years group, only the subfoveal choroidal thickness can be used for diagnosis.

The nasal choroidal thickness is not able to differentiate eyes with MDCA from those with PDCA. Instead, the choroidal thickness cut-off value of 62 μm at the subfovea (sensitivity, 71%; specificity, 72%), 73 μm at the temporal location (sensitivity, 67%; specificity, 90%), 83 μm at the superior location (sensitivity, 67%; specificity, 80%), 84.5 μm at the inferior location (sensitivity, 81%; specificity, 65%) can be used to define the eyes with MDCA.7 Although the area under curve of subfoveal choroidal thickness is not the largest among all the locations, the cut-off value is still based on the subfoveal choroidal thickness, because the measurement at the fovea is easier to take and more accurate than the parafoveal points.

Summary of OCT-Based Classification of Myopic Maculopathy. Combining the hallmarks of myopic maculopathy in pathologic myopia, a new classification based on OCT findings has been proposed. In this new system, diffuse choroidal atrophy, for PDCA and MDCA, is suggested to be called “peripapillary choroidal thinning” and “macular choroidal thinning” (Table 3). Based on these results, the cut-off value of choroidal thickness as a diagnostic tool for diffuse atrophy has been added to this system. That is, peripapillary choroidal thinning is defined as a choroidal thickness of less than 56.5 μm at 3000 μm nasal from the fovea and macular choroidal thinning is defined as a choroidal thickness of less than 62 μm at the subfovea. Patchy atrophy and MNV-related macular atrophy are not simply owing to atrophy, but to holes in Bruch's membrane. Patchy atrophy is seen as a well-defined, grayish white lesion by fundus photograph seldomly involving the central fovea which is appropriate called as “extrafoveal Bruch's membrane defects” by OCT definition. In contrast, foveal Bruch's membrane defects is named for macular atrophy, that is, category 4 in the META-PM classification, for both MNV-related and patchy-related disease. In addition, myopic traction maculopathy and dome-shaped macula both as potential vision-threatening macular complications that only can be detected on OCT are also included in the OCT-based classification of myopic maculopathy.

Further studies are needed to validate whether the cut-off values of choroidal thickness from a single high myopia clinic can work well in clinical practice. More studies should investigate the role of the choroid in high myopia, not only by a measurement of the thickness, but also by examining other parameters, such as choroidal blood flow and the morphologic and vascular features of the choroid. Longitudinal studies on choroidal change will help us to depict a real-world picture of the choroidal changes in patients with high myopia.

Future Perspectives

Diagnostic methods as well as therapies have been greatly advancing for pathologic myopia. In particular, anti-VEGF therapies as well as vitreoretinal surgeries have improved the anatomical and visual outcomes of myopic MNV and myopic traction maculopathy. However, therapeutic approaches are not sufficient for other complications, such as choroidal atrophies and optic nerve damage. The diagnosis of glaucoma is difficult in eyes with pathologic myopia, because of a deformity of the optic disc appearance and the scotoma caused by myopic macular lesions and a large gamma zone. How to diagnose glaucoma in its early stage of patients with pathologic myopia needs to be established in the future, because the optic nerve damage is considered an important cause of blindness in pathologic myopia.

Posterior staphyloma causes a deformity of the area containing visually important tissues, namely, macula and the optic nerve. It is in contrast with most of the axial elongation in myopic eyes in general occurs in the equator of the eye. Thus, a formation of posterior staphyloma mechanically damages these nerve tissues. Before developing irreversible blinding complications, approaches to prevent and treat staphylomas are expected.

Finally, clarifying which genes are responsible for pathologic myopia (not high myopia) is important. With this information, it will finally become clear whether or not pathologic myopia is different from myopia in general. In addition, identifying children who will become pathologically myopic will be possible.

Based on the consistent definition of pathologic myopia and the objective detection of its major feature—posterior staphyloma—future studies regarding the global prevalence of pathologic myopia and its impact on vision and quality of life294 should be undertaken.

Acknowledgments

The authors thank Monica Jong for facilitation of the process.

Supported by the International Myopia Institute. The publication costs of the International Myopia Institute reports were supported by donations from the Brien Holden Vision Institute, Carl Zeiss Vision, CooperVision, Essilor, and Alcon.

Disclosure: K. Ohno-Matsui, Santen (C), Nevakar (C); P.-C. Wu, None; K. Yamashiro, None; K. Vutipongsatorn, None; Y. Fang, None; C.M.G. Cheung, Novartis, Bayer, Roche, Topcon, Zeiss, Allergan, and Boehringer Ingelheim (F, C); T.Y.Y. Lai, Bayer Healthcare (C, F, R), Boehringer Ingelheim (C), Roche (C, F, R), Novartis (C, F, R); Y. Ikuno, Novartis Pharma (R); S.Y. Cohen, Allergan (C), Bayer (C), Novartis (C), Thea (C), Roche (C); A. Gaudric, None; J.B. Jonas, Europäische Patentanmeldung 16 720 043.5 and Patent application US 2019 0085065 A1 „Agents for use in the therapeutic or prophylactic treatment of myopia or hyperopia“ (P)

References

41. Kiefer AK, Tung JY, Do CB, et al. Genome-wide analysis points to roles for extracellular matrix remodeling, the


231. Cai B, Yang J, Li S, et al. Comparison of the efficacy of intravitreal ranibizumab for choroidal neovascularization...


Fledelius HC, Jacobsen N, Li XQ, Goldschmidt E. Choroidal thickness at age 66 years in the Danish high myopia study cohort 1948 compared with follow-up data on visual acuity over 40 years: a clinical update adding spectral domain optical coherence tomography. *Acta Ophthalmol.* 2018;96:46–50.


